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CHEMICAL ENGINEERING

VOLUME 1

SIXTH EDITION

Fluid Flow, Heat Transfer and Mass Transfer

J. M. COULSON

Late Emeritus Professor of Chemical Engineering University of Newcastle-upon-Tyne

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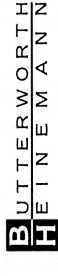
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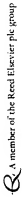
WITH

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Contents

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Prof	essor	Professor J. M. Coulson	ulson	xiii
Pref	ace u	Preface to Sixth Edition	dition	A.Y
Pref	ace u	Preface to Fifth Edition	dition	xvii
Pref	ace u	Preface to Fourth Edition	Edition	xix
Pref	ace u	Preface to Third Edition	dition	xxi
Pref	ace u	Preface to Second Edition	Edition	xxiii
Pref	ace ta	Preface to First Edition	lition	XX
4ck)	эрмон	Acknowledgements		xxvii
	Ë	S and D	Units and Dimensions	-
	-			•
	<u> </u>	Systems of units	ni f units	- 7
			The centimetre-gram-second (egs) system	C+ -
			the metre-kitogram - second (mks system) and the Système International d'Unités (SI)	7
		1.23	The foot-pound-second (fps) system The British enomerating system	ws w
			Non-coherent	n e
			system employing pound mass and pound force simultaneously	,
		0.7.1	Derived ands Thermal theorie	Φ (-
			Molar units	- 90
		1.2.9	Electrical units	· ж
	2:	Conversion	Zonversian of units	3
	च् <u>र</u> ;	Dimensio	Dimensional analysis	2
	7 Y	Buckingh	Suckingham's II (negreen) Sectoficition of the female and ensec dimensions	2 5
	?	1.6.1	Vector and scalar quantities	200
		1.6.2	Quantity mass and inertia mass	7
	1.3	Further reading	guipa	23
	33	References		32
	1.9	Nomenclature	in c	Ħ
Part	G	Part 1 Fluid Flow		ž
	•			3
ri	Flox	v of Flui	Flow of Fluids Energy and Momentum Relationships	27
	7.7	Introduction Internal energy	33 PETEV	22
				ì

CONTENTS	4.4. Converging -diverging nozzles for gas flow 4.4.1 Maximum flow and critical pressure ratio 4.4.2 The pressure and area for flow 4.4.3 Effect of back-pressure on flow in nozzle 4.5.1 Enerty balance for flow of ideal eas	4.5.2 Isothermal flow of an ideal gas in a horizontal pipe 4.5.3 Non-isothermal flow of an ideal gas in a horizontal pipe 4.5.4 Adiabatic flow of an ideal gas in a horizontal pipe 4.5.5 Flow of non-ideal gases 4.6 Shock waves	4.7 Further reading 178 4.5 References 179 4.9 Nomenclature 179	5. Flow of Multiphase Mixtures 5.1 Introduction 5.2 Two-phase gas (vapour)-liquid flow 5.2.1 Introduction	5.2.2 Flow regimes and flow patterns 5.2.3 Hold-up 5.2.4 Pressure, momentum, and energy relations 5.2.5 Freezing	5.3 Flow of 5.3.1 S.3.1 S.3.1 S.3.2	5.3.4 Coarse solids 5.3.4 Coarse solids in horizontal flow 5.3.5 Coarse solids in vertical flow 5.4 Flow of gas-solids mixtures 5.4 I Coarsell considerations	5.4.2 Horizonta transport 5.4.3 Vertical transport 5.4.4 Practical applications 5.5 Futher reading 5.6 References	57 6. Flow	6.1 Introduction 6.2 Fluid pressure 6.2.1 Static pressure 6.2.2 Pressure measuring devices 6.2.3 Pressure signal transmission — the differential pressure cell 6.2.4 Intelligent pressure transmitters	6.2.5 6.3.1 6.3.1 6.3.2 6.3.3 6.3.3 6.3.3	6.3.5 The ventum meter 6.3.6 Pressure recovery in ordice-type meters 6.3.7 Variable area meters—roamneters	6.3.8 The notch or weir 6.3.9 Other methods of measuring flowrates 6.4 Further reading 6.5 References 6.6 Nomenclature
CONTENTS	of fluid The incompressible fluid (liquid) The ideal gas The non-ideal gas	2.4.1 Continuity 2.4.2 Momentum changes in a fluid 2.4.3 Integry of a fluid in motion 2.4.4 Pressure and fluid head 2.4.4 Pressure and fluid head 4.5 Constant flow per unit area 2.4.6 Constant flow per unit area	ure volume relationships Incompressible fluids Compressible fluids ional or vortex motion in a fluid		×	3.1 Introduction 3.2 The nature of fluid flow 3.2.1 Plow over a surface 60	Flow in a pipe onin the state of a Newtonian fluid Shearing characteristics of a Newtonian fluid Pressure drop for flow of Newtonian liquids through a pipe	Reynolds number and shear stress Velocity distributions and volumetric flowrates for streamline flow The transition from laminar to turbulent flow in a pipe Velocity distributions and volumetric flowrates for includent flow Flow through curved pipes	s in behaviour	Time-dependent behaviour Viscoelastic behaviour Viscoelastic behaviour Characterisation of non-Newtonian fluids Dimensionless characterisation of viscoelastic flows Relation between rhealogy and structure of material Kreaniine flow in ringes and channels of regular geometry	Turbulent flow The transition from laminar to turbulent flow rer reading rences	4. Flow of Compressible Fluids 143	4.1 Introduction 4.2 Flow of gas through a nozzle or orifice 4.2.1 Isothermal flow 4.2.2 Non-isothermal flow 4.2.2 Non-isothermal flow

Scale-up of stirred vessels Turbulem mixing

27

Mixing mechanisms

5

7.4.1 Low viscosity systems 7.4.2 High viscosity systems Flow patterns in stirred tanks Rate and time for mixing

Mixing equipment

7.5

7.7.4 Static mixers
7.7.5 Other types of mixer
Mixing in continuous systems
Further reading

Introduction—types of mixing 7.1.1 Single-phase liquid mixing 7.1.2 Mixing of immixible liquids

Liquid Mixing

≅

7

13 10 10 10 10 10 10 10			in all		
sing the legack by the legack			e d	Hear	weight har machinism
a viniting 274 9.33 a viniting 275 9.34 b the liquids 275 9.4 Heat unit for pumping through pipelines of beat transfer 8 9.1 a viniting 275 9.4 Heat unit for pumping through pipelines of beat transfer 8 9.1 a viniting 275 9.4 Heat unit for pumping through pipelines of beat transfer 8 9.1 a viniting 275 9.4 Heat unit for pumping through pipelines 9 9.1 a viniting 275 9.4 Heat unit for pumping through pipelines 9 9.1 a viniting 312 9.7 9.4 a viniting 312 9.7 Bolding 9.8 a viniting 275 9.8 a viniting 312 9.7 Bolding 9.8 a viniting 312 9.7 Bolding 9.8 a viniting 275 9.8 a viniting 312 9.7 Bolding 9.8 a viniting 312 9.7 a viniting 322 9.7 a viniting 3				23.1	Conduction through a plane wall
d mixing			ut. B	9,3,2	Thermal resistances in series
Dec Principal Color	and the second s		:	93.3	Conduction through a thick-walled tube
one 275 94 44 Heat transportations strig applications 277 94 94,3 strig applications 277 94 94,3 strig applications 277 94 94,3 red vessels 280 94,2 94,3 red vessels 280 94,2 94,3 red vessels 280 94,2 94,3 series 282 94,2 94,3 series 283 283 95,4 series 284 95,2 94,3 series 381 381 95,3 series 310 96,1 96,2 series 311 97,2 94,3 series 311 97,3 94,3 series 314 97,3 94,3 series 314 97,3 94,3 series 344 97,4 97,2 series 344 97,4 97,1 series <t< td=""><td></td><td></td><td></td><td>9.3,4</td><td>Conduction through a spherical shell and to a particle</td></t<>				9.3,4	Conduction through a spherical shell and to a particle
ting applications 177 9.4 4.1 6.4 1				9,3,5	Unsteady state conduction
sing applications	• • •			9.3.6	Conduction with internal heat source
ing applications 276 9.44 1.7				Heat	innister by convection
red vessels 277 277 24.3 red vessels 280 24.4 stems 281 282 24.4 stems 282 282 24.4 stems 282 283 24.4 stems 283 294 26.2 stems 283 294 26.2 stems 284 26.2 stems 284 26.2 stems 284 26.2 stems 285 29.4 stems 285 29.4 stems 286 26.2 stems 286				7	Salufat and forced convection
rect vessels 227 24, 24, 24, 24, 24, 24, 24, 24, 24, 24,	The state of the s	***		2 4 2	Capacidation of University and State Consection
red vessels 257 269 9.4.5 acrost stems 258 259 4.7 acrost stems 250 5.5	iii			9.4.4	Forced convertion outside tubes
terms verms ve		, J.		9.4.5	Plaw in non-circular sections
stems				9.4.6	Convection to spherical particles
sterins sterin				9.4.7	Natural convection
Secondary			9.5	Heat	transfer by radiation
1985 1985	\$	9 5		9.5.1	Introduction
tion increments	nks	20		9.5.3	Radiation from a black body
1975 1975				9.5.3	Radiation from real surfaces
10				9.5.4	Radiation transfer between black surfaces
10		× ×		9.5.5	Radiation transfer between grey surfaces
197 197					Kadianon Irom gases
130 96.2 131 9.7 Boling 131 9.7 131 9.7 Boling 131 9.7		bot.		_ `	transfer in the condensation of valvants
10				9.0.1	Candengorism on venical and memoral subsections.
11 9.6.4 12 13 13 14 9.7 13 14 9.7 14 9.7 15 15 15 15 16 17 17 18 18 18 18 18 18 18				9.63	Property condensation
114 2.7 Bolling 2.7		***		9.6.4	Condensation of mixed vapours
314 97.1		***			lg liquids
114 9.7.2 115 9.7.5 116 9.7.5 117 9.7.5 118 9.7.5 119 9.7.5 119 9.7.5 119 9.7.5 120 9.7.5 121 9.7.5				9.7.1	Conditions for boiling
State Stat				9.7.2	Types of Dotting
State Stat		****		2.4.0	Acadesis based on bubble champeristics
State Stat				9.7.5	Sub-cooled boiling
Section Sect	aipment for liquids	w/w/ ***		9.7.6	
displacement rotary pumps displacement rotary pumps nent for gases not rotary compressors fugal and turbocompressors fugal and unbocompressors required for the compression of gases special for the compression of		man.			transfer in reaction vessels
nutringal pump and runty compressors and runty compressors and unbocompressors appears and in for pumping and unbocompressors appears and in for pumping through pipelines and unbocompressors appears and in for pumping through pipelines and unbocompressors and in for pumping through pipelines and unbocompressors and in for pumping through pipelines and unbocompressors and in for pumping through pipelines and overall evertheients of heat transfer and overall evertheients and overall evertheients and overall evertheients of heat transfer and overall evertheients and overall evertheients of heat transfer and overall evertheients of heat transfer and overall evertheients and overal	wary pumps			9.8.1	Helical cooling coils
1999 1999		****		9,50	Jacketed vessels
August A	Service Control of the Control of th	7	00		lime required for nealing of cooling
147 147 149		99	6.6		Conservation description
required for the compression of gases 358 9.9.3 158 158 9.9.4 158 159.5 158 159.5 158 159.5 158 159.5 159 159.5 150 150 159.5 150 150 150 150 150 150 150 150 1	JO.	47		9.0.2	Basic components
1984 1994 1994 1995				9.9.3	Mean temperature difference in multipass exchangers
25.9				766	Film coefficients
1996 1996 1996 1996 1996 1996 1996 1996 1997		44.5	. "	9.9.5	
ments for pumping through pipelines (1988) (1988) (1993) (9.6.6	
374 9.10 Other for 510 376 9.10 Other for 510 376 9.10 Other for 510 376 9.10 376 9.10 377 9.10 377 9.10 379 9.10 3.10 3.10 3.10 3.10 3.10 3.10 3.10 3				9.9.7	
376 9.10 Other to 376 9.10 Other to 9.10		***		-	Transfer units
376 2.10.2 377 3.10.2 377 9.10.3 9.10.3 9.10.4 9.10.4 9.10.4 9.11.7 9.11.1 381 9.11.2 9.11.2 9.11.3 9.11.3 9.12.7 9.11.3 9.13.8 9.13.8 9.14.8 9.15.8 9.14.8 9.15.9 9.14.10.8 9.15.10.8	Gases				₫ .
377 9,10.3 379 9,10.4 379 9,11 Thermal 9,11 Thermal 381 9,11 Thermal 381 9,11 Thermal 9,11 3,11,3 14 9,11 Thermal 9,11 Temperature difference 384 9,12 Further 7 September 9,11 Nomenci	automa Lindon	100	15.4	0.50	
379 9.10.4	nces.		il unti-	9.103	
381 9.10.5 179 4 9.10.5 181 9.11.1 181 9.11.3 181 9.11.3 181 9.12 Further i further is of heat transfer 381 9.13 Reference				9.10.4	
381 9.11.1 1100 11.12 11.13	ınsfer	(to a late			1
381 9,11.2 381 9,11.3 4,11.3 51.1 51.1 52.1 53.1 53.1 53.1 53.1 54.1 55.1 56.1 56.1 56.1 56.1 56.1 56.1 56		*****		_	7
381 9.12 Further and overall exefficients of heat transfer 381 9.13 Reference 384 9.14 Nomenet				9.11.2	
nd-overall exefficients of heat transfer 58.1 38.1 9.1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2 1.2	uction	70 Y		.	
384	CONSIDERATIONS CONSIDERATIONS CONTRACTOR TO THE PROPERTY OF TH				er reading
	Individual and Overal exements of the management	38.1	**************************************		nclature
			·		

8,88,80 7,80,80 9,90

Part 2 Heat Transfer

9. Heat Transfer

8.3

Introduction

Pumping of Fluids

∞;

Nomenclature

References

7.8 7.9 7.10

46.5

Marian Company

CONTENTS	11.6 The boundary layer for heat transfer		surface temperature 11.64 Heat transfer for streamline flow over a plane surface — constant	11.7 The boun	11.8 Further reading	ger (gr.)	12. Momentum, Heat, and Mass Transfer	12.1	12.2 Transfer by molecular diffusion 12.2.1 Momentum transfer		12.2.3 Mass transfer			12.3.1 The nature of turbulent flow	12.3.2	12.4 Universal velocity profile	12,4,2	12.4.3	1.4.4.5.		12.4.7	12,4,8	12.5	12.6		12.8.1	12.8.3 Taylor—Prandtl modification of Reynolds analogy for heat	-	12.8.4 Use of universal velocity profile in Reynolds analogy	-	12.9 Further retaining 12.10 References	12.11 Nomenclature		13. Humidiheation and Water Cooling	13.1 Introduction	13.2.1	13.2.2 Wet-bulb temperature	***	13.3.2 Enthalpy—humidity chart
	201	573	£ 8 8 8	576	S81	288 288		593	593	, s	597	5550 KID	602	614	819	819				169			ung f		940	£	651	150	655	959		100	663		\$99	675	675 677	- F 3	
CONTENTS	Part 3 Mass Transfer	10. Mass Transfer	2 0	10.2.1 Properties of binary mixtures 10.2.2 Equimolecular counterdiffusion		10.2.6 General case for gas-phase mass transfer		10.2.9 Unsteady-state mass transfer	10.3.1	10.3.2 Maxwell's law of diffusion			10.5.1 The two-titut orders	The film - penetration theory	10.5.4 Mass transfer to a sphere in a homogenous fluid			10.5.8 Countercurrent mass transfer and transfer units 10.6. Mass transfer and chemical transfer			10.7 Mass transfer and chemical reaction in a catalyst petied in 2.1. Fine sharefore			10,7,4 Mass transfer and chemical reaction with a mass transfer resistance external to the pellet	Il studies of mass transfer	10.8.1 The <i>j</i> -factor of Criticon and Colourn for now in macs	Effect of surface roughness and form drag	10.8.4 Mass transfer from a fluid to the surface of particles		10.11 Nomenclature	The state of the s	Fart 4 Nomembum, fical and Mass Transfer	11. The Boundary Laver	Introvbiction			11.4.1 The turbulent partion	11.5 Boundary layer theory applied to pipe flow	11.5.1 Entry conditions 11.5.2 Application of the boundary-layer theory

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-
•
7
ш
-
~
=
ਨ
ರ
ၓ
ၓ
ŏ
ၓ

:₹

13.4	Determination of humidity
13.5	Humidification and dehumidification
	13.5.1 Methods of increasing humidity
	13.5.2 Dehumidification
13.6	Water cooling
	13.6.1 Cooling towers
	13.6.2 Design of natural-draught towers
	13.6.3 Height of packing for both natural and mechanical draught to
	13.6.4 Change in air condition
	13.6.5 Temperature and humidity gradients in a water cooling tower
	13,6.6 Evaluation of heat and mass transfer coefficients
	13.6.7 Humidifying towers
13,7	Systems other than air-water
13.8	Further sending
13.9	References
13.10	Nomenclature
Appendix	
AI. T	A1. Tables of physical properties
A2. S	A2. Steam tables
A3. N Folds	A3. Mathematical tables Fold-out charts
Problems	

Index

Professor J. M. Coulson

echanical draught towers

single-handed, a very demanding assignment. During this period he collaborated with Sir on "The Manufacture of Nitrotoluene". He published research papers on heat transfer JOHN COULSON, who died on 6 January 1990 at the age of 79, came from a family with close involvement with education. Both he and his twin brother Charles (renowned physicist and mathematician), who predeceased him, became professors, John did his undergraduate studies at Cambridge and then moved to Imperial College where he took the postgraduate course in chemical engineering -the normal way to qualify at that time - and then carried out research on the flow of fluids through packed beds. He then became an Assistant Lecturer at Imperial College and, after war-time service in the Royal Ordnance Factories, returned as Lecturer and was subsequently promoted to a Readership. At Imperial College he initially had to run the final year of the undergraduate course almost Frederick (Ned) Warner to write a model design exercise for the I. Chem. E. Home Paper and evaporation, on distillation, and on liquid extraction, and co-authored this textbook of Chemical Engineering. He did valiant work for the Institution of Chemical Engineers which awarded him its Davis medal in 1973, and was also a member of the Advisory Board for what was then a new Pergamon journal, Chemical Engineering Science.

- In 1954 he was appointed to the newly established Chair at Newcastle-upon-Tyne, where Chemical Engineering became a separate Department and independent of Mechanical He took a period of secondment to Heriot Watt University where, following the splitting of the joint Department of Chemical Engineering with Edinburgh, he acted as adviser and de facto Head of Department. The Scottish university awarded him an Honorary D.Sc. in Engineering of which it was formerly part, and remained there until his retirement in 1975.

John's first wife Dora sadly died in 1961; they had two sons, Anthony and Simon. He remarried in 1965 and is survived by Christine.

JFR

CHAPTER 9

Heat Transfer

9.1. INTRODUCTION

In the majority of chemical processes heat is either given out or absorbed, and fluids must often be either heated or cooled in a wide range of plant, such as furnaces, evaporators, distillation units, dryers, and reaction vessels where one of the major problems is that of transferring heat at the desired rate. In addition, it may be necessary to prevent the loss of heat from a hot vessel or pipe system. The control of the flow of heat at the desired rate forms one of the most important areas of chemical engineering. Provided that a temperature difference exists between two parts of a system, heat transfer will take place in one or more of three different ways.

Conduction. In a solid, the flow of heat by conduction is the result of the transfer of vibrational energy from one molecule to another, and in fluids it occurs in addition as a result of the transfer of kinetic energy. Heat transfer by conduction may also arise from the movement of free electrons, a process which is particularly important with metals and accounts for their high thermal conductivities.

Convection. Heat transfer by convection arises from the mixing of elements of fluid. If this mixing occurs as a result of density differences as, for example, when a pool of liquid is heated from below, the process is known as natural convection. If the mixing results from eddy movement in the fluid, for example when a fluid flows through a pipe heated on the outside, it is called forced convection. It is important to note that convection requires mixing of fluid elements, and is not governed by temperature difference alone as is the case in conduction and radiation.

Radiation. All materials radiate thermal energy in the form of electromagnetic waves. When this radiation falls on a second body it may be partially reflected, transmitted, or absorbed. It is only the fraction that is absorbed that appears as heat in the body.

9.2. BASIC CONSIDERATIONS

9.2.1. Individual and overall coefficients of heat transfer

In many of the applications of heat transfer in process plants, one or more of the mechanisms of heat transfer may be involved. In the majority of heat exchangers heat passes through a series of different intervening layers before reaching the second fluid (Figure 9.1). These layers may be of different thicknesses and of different thermal conductivities. The problem of transferring heat to crude oil in the primary furnace before it enters the first distillation column may be considered as an example. The heat from the flames passes by radiation and convection to the pipes in the furnace, by conduction through the

Figure 9.1. Heat transfer through a composite wall

pipe walls, and by forced convection from the inside of the pipe to the oil. Here all three modes of transfer are involved. After prolonged usage, solid deposits may form on both the inner and outer walls of the pipes, and these will then contribute additional resistance to the transfer of heat. The simplest form of equation which represents this heat transfer operation may be written as:

$$Q = UA\Delta T \tag{9.1}$$

where Q is the heat transferred per unit time, A the area available for the flow of heat, $\Delta \Gamma$ the difference in temperature between the flame and the boiling oil, and U is known as the overall heat transfer coefficient (W/m² K in SI units).

At first sight, equation 9.1 implies that the relationship between Q and ΔT is linear. Whereas this is approximately so over limited ranges of temperature difference for which U is nearly constant, in practice U may well be influenced both by the temperature difference and by the absolute value of the temperatures.

If it is required to know the area needed for the transfer of heat at a specified rate, the temperature difference ΔT , and the value of the overall heat-transfer coefficient must be known. Thus the calculation of the value of U is a key requirement in any design problem in which heating or cooling is involved. A large part of the study of heat transfer is therefore devoted to the evaluation of this coefficient.

The value of the coefficient will depend on the mechanism by which heat is transferred, on the fluid dynamics of both the heated and the cooled fluids, on the properties of the materials through which the heat must pass, and on the geometry of the fluid paths. In solids, heat is normally transferred by conduction; some materials such as metals have a high thermal conductivity, whilst others such as ceramics have a low conductivity. Transparent solids like glass also transmit radiant energy particularly in the visible partiof the spectrum.

Liquids also transmit heat readily by conduction, though circulating currents are frequently set up and the resulting convective transfer may be considerably greater than the transfer by conduction. Many liquids also transmit radiant energy. Gases are poor conductors of heat and circulating currents are difficult to suppress; convection is therefore much more important than conduction in a gas. Radiant energy is transmitted with only limited absorption in gases and, of course, without any absorption in vacuo. Radiation is the only mode of heat transfer which does not require the presence of an intervening

If the heat is being transmitted through a number of media in series, the overall heat transfer coefficient may be broken down into individual coefficients h each relating to a single medium. This is as shown in Figure 9.1. It is assumed that there is good contact between each pair of elements so that the temperature is the same on the two sides of

If heat is being transferred through three media, each of area A, and individual coefficients for each of the media are h_1 , h_2 , and h_3 , and the corresponding temperature changes are ΔT_1 , ΔT_2 , and ΔT_3 then, provided that there is no accumulation of heat in the media, the heat transfer rate Q will be the same through each. Three equations, analogous to equation 9.1 can therefore be written:

$$Q = h_1 A \Delta T_1$$

$$Q = h_2 A \Delta T_2$$

$$Q = h_3 A \Delta T_3$$

$$A T_1 = \frac{Q}{A} \frac{1}{h_1}$$

$$\Delta T_2 = \frac{Q}{A} \frac{1}{h_2}$$

$$\Delta T_3 = \frac{Q}{A} \frac{1}{h_3}$$

$$\Delta T_3 = \frac{Q}{A} \frac{1}{h_3}$$

$$\Delta T_3 = \frac{Q}{A} \frac{1}{h_3}$$

$$(9.3)$$
Adding:
$$\Delta T_1 + \Delta T_2 + \Delta T_3 = \frac{Q}{A} \left(\frac{1}{h_1} + \frac{1}{h_2} + \frac{1}{h_3}\right)$$

Noting that $(\Delta T_1 + \Delta T_2 + \Delta T_3) = \text{total temperature difference } \Delta T$:

$$\Delta T = \frac{Q}{A} \left(\frac{1}{h_1} + \frac{1}{h_2} + \frac{1}{h_3} \right) \tag{9.4}$$

From equation 9.1:
$$\Delta T = \frac{Q}{A} \frac{1}{U}$$
 (9.5)

Comparing equations 9.4 and 9.5:

$$\frac{1}{U} = \frac{1}{h_1} + \frac{1}{h_2} + \frac{1}{h_3} \tag{9.6}$$

The reciprocals of the heat transfer coefficients are resistances, and equation 9.6 therefore illustrates that the resistances are additive.

In some cases, particularly for the radial flow of heat through a thick pipe wall or cylinder, the area for heat transfer is a function of position. Thus the area for transfer applicable to each of the three media could differ and may be A_1 , A_2 and A_3 . Equation 9.3 then becomes:

$$\Delta T_1 + \Delta T_2 + \Delta T_3 = Q\left(\frac{1}{h_1 A_1} + \frac{1}{h_2 A_2} + \frac{1}{h_3 A_3}\right) \tag{9.7}$$

Equation 9.7 must then be written in terms of one of the area terms A_1 , A_2 , and A_3 , or sometimes in terms of a mean area. Since Q and ΔT must be independent of the particular

382

area considered, the value of U will vary according to which area is used as the basis. Thus equation 9.7 may be written, for example:

$$Q = U_1 A_1 \Delta T$$
 or $\Delta T = \frac{Q}{U_1 A_1}$

This will then give U_1 as:

$$\frac{1}{U_1} = \frac{1}{h_1} + \frac{A_1}{A_2} \left(\frac{1}{h_2} \right) + \frac{A_1}{A_3} \left(\frac{1}{h_3} \right) \tag{9.8}$$

In this analysis it is assumed that the heat flowing per unit time through each of the media

the same.

Now that the overall coefficient U has been broken down into its component parts, each of the individual coefficients h_1 , h_2 , and h_3 must be evaluated. This can be done from a knowledge of the nature of the heat transfer process in each of the media. A study will therefore be made of how these individual coefficients can be calculated for conduction, convection, and radiation.

9.2.2. Mean temperature difference

Where heat is being transferred from one fluid to a second fluid through the wall of a vessel and the temperature is the same throughout the bulk of each of the fluids, there is no difficulty in specifying the overall temperature difference ΔT . Frequently, however, each fluid is flowing through a heat exchanger such as a pipe or a series of pipes in parallel, and its temperature changes as it flows, and consequently the temperature difference is continuously changing. If the two fluids are flowing in the same direction (co-current flow), the temperatures of the two streams progressively approach one another as shown in Figure 9.2. In these circumstances the outlet temperature of the heating fluid must always be higher than that of the cooling fluid. If the fluids are flowing in opposite directions (countercurrent flow), the temperature difference will show less variation throughout the heat exchanger as shown in Figure 9.3. In this case it is possible for the cooling liquid to leave at a higher temperature than the heating liquid, and one of the great advantages of

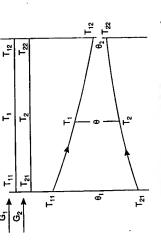


Figure 9.2. Mean temperature difference for co-current flow

HEAT TRANSFER

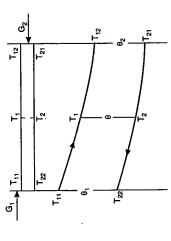


Figure 9.3. Mean temperature difference for countercurrent flow

countercurrent flow is that it is possible to extract a higher proportion of the heat content of the heating fluid. The calculation of the appropriate value of the temperature difference for co-current and for countercurrent flow is now considered. It is assumed that the overall heat transfer coefficient *U* remains constant throughout the heat exchanger.

It is necessary to find the average value of the temperature difference θ_m to be used in the general equation:

$$Q = UA\theta_m (equation 9.1)$$

Figure 9.3 shows the temperature conditions for the fluids flowing in opposite directions, a condition known as countercurrent flow.

The outside stream specific heat C_{p1} and mass flow rate G_1 falls in temperature from T_{11} to T_{12}

The inside stream specific heat C_{p2} and mass flow rate G_2 rises in temperature from T_{21} to T_{22} . Over a small element of area dA where the temperatures of the streams are T_1 and T_2 .

$$\theta = T_1 - T_2$$

$$d\theta = dT_1 - dT_2$$

The temperature difference:

Heat given out by the hot stream $= dQ = -G_1C_{p1} dT_1$

Heat taken up by the cold stream $= dQ = G_2C_{p2} dT_2$

$$d\theta = -\frac{dQ}{G_1C_{p1}} - \frac{dQ}{G_2C_{p2}} = -dQ\left(\frac{G_1C_{p1} + G_2C_{p2}}{G_1C_{p1} \times G_2C_{p2}}\right) = -\psi \,dQ \quad \text{(say)}$$

 $\theta_1 - \theta_2 = \psi Q$

Over this element: $U dA\theta = dQ$

$$U \, dA\theta = -\frac{d\theta}{\psi}$$

If U may be taken as constant:

$$-\psi U \int_0^A dA = \int_{\theta_1}^{\theta_2} \frac{d\theta}{\theta}$$
$$-\psi U A = -\ln \frac{\theta_1}{\theta_2}$$

From the definition of θ_m , $Q = UA\theta_m$.

$$\theta_1 - \theta_2 = \psi Q = \psi U A \theta_m = \ln \frac{\theta_1}{\theta_2} (\theta_m)$$

$$\theta_m = \frac{\theta_1 - \theta_2}{\ln(\theta_1/\theta_2)}$$

and:

(6.6)

where θ_m is known as the logarithmic mean temperature difference.

UNDERWOOD⁽¹⁾ proposed the following approximation for the logarithmic mean temperature difference:

$$(\theta_m)^{1/3} = \frac{1}{2}(\theta_1^{1/3} + \theta_2^{1/3})$$
 (9.10)

and the logarithmic mean values coincide at 39 K.

If the two fluids flow in the same direction on each side of a tube, co-current flow is and, for example, when $\theta_1 = 1 \text{ K}$ and $\theta_2 = 100 \text{ K}$, θ_m is 22.4 K compared with a logarithmic mean of 21.5 K. When $\theta_1 = 10$ K and $\theta_2 = 100$ K, both the approximation

taking place and the general shape of the temperature profile along the tube is as shown in Figure 9.2. A similar analysis will show that this gives the same expression for θ_m , the logarithmic mean temperature difference. For the same terminal temperatures it is important to note that the value of θ_m for countercurrent flow is appreciably greater than the value for co-current flow. This is seen from the temperature profiles, where with co-current flow the cold fluid cannot be heated to a higher temperature than the exit temperature of the hot fluid as illustrated in Example 9.1.

Example 9.1

at 300 K. If the overall coefficient of heat transfer is constant at 2 kW/m²K, calculate the surface area required in (a) a countercurrent concentric tube exchanger, and (b) a co-current flow concentric tube exchanger. A heat exchanger is required to cool 20 kg/s of water from 360 K to 340 K by means of 25 kg/s water entering

Solution

Heat load: $Q = 20 \times 4.18(360 - 340) = 1672 \text{ kW}$

The cooling water outlet temperature is given by:

$$1672 = 25 \times 4.18(\theta_2 - 300)$$
 or $\theta_2 = 316 \text{ K}$

(a) Counterflow

In equation 9.9:

$$\theta_m = \frac{44 - 40}{\ln(44/40)} = 41.9 \text{ K}$$

HEAT TRANSFER

$$= \frac{1672}{2 \times 41.9}$$
$$= \frac{19.95 \text{ m}^2}{2 \times 41.9}$$

$$\theta_m = \frac{60 - 24}{\ln(60/24)} = 39.3 \text{ K}$$

In equation 9.9:

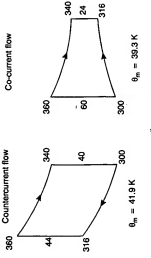
Heat transfer area:

$$A = \frac{1672}{2 \times 39.3}$$

$$A = \frac{1672}{2 \times 39.3}$$

$$= \frac{21.27 \text{ m}^2}{3 \times 39.3}$$

It may be noted that using Underwood's approximation (equation 9.10), the calculated values for the mean temperature driving forces are 41.9 K and 39.3 K for counter- and co-current flow respectively, which agree exactly with the logarithmic mean values.



24

Figure 9.4. Data for Example 9.1

9.3. HEAT TRANSFER BY CONDUCTION

9.3.1. Conduction through a plane wall

This important mechanism of heat transfer is now considered in more detail for the flow of heat through a plane wall of thickness x as shown in Figure 9.5.

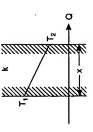


Figure 9.5. Conduction of heat through a plane wall

The rate of heat flow Q over the area A and a small distance dx may be written as:

$$Q = -kA \left(\frac{dT}{dx}\right) \tag{9.11}$$

which is often known as Fourier's equation, where the negative sign indicates that the temperature gradient is in the opposite direction to the flow of heat and k is the thermal conductivity of the material. Integrating for a wall of thickness x with boundary temperatures T_1 and T_2 , as shown in Figure 9.5:

$$Q = \frac{kA(T_1 - T_2)}{r} \tag{9.12}$$

Thermal conductivity is a function of temperature and experimental data may often be expressed by a linear relationship of the form:

(9.13)where k is the thermal conductivity at the temperature T and k_0 and k' are constants. Combining equations 9.11 and 9.13: $k = k_0(1 + k'T)$

$$-k \, dT = -k_0 (1 + k'T) dT = \frac{Q \, dx}{A}$$

Integrating between the temperature limits
$$T_1$$
 and T_2 ,
$$-\int_{T_1}^{T_2} k \, dT = (T_1 - T_2)k_0 \left\{ 1 + k' \left(\frac{T_1 + T_2}{2} \right) \right\} = Q \int_{x_1}^{x_2} \frac{dx}{A}$$
 (9.14)

Where k is a linear function of T, the following equation may therefore be used:

$$k_a(T_1 - T_2) = Q \int_{x_1}^{x_2} \frac{dx}{A}$$
 (9.15)

where k_a is the arithmetic mean of k_1 and k_2 at T_1 and T_2 respectively or the thermal Where k is a non-linear function of T, some mean value, k_m will apply, where: conductivity at the arithmetic mean of T_1 and T_2 .

increase in molecular mass and increases with the temperature. In addition, for gases the From Table 9.1 it will be seen that metals have very high thermal conductivities, nonhas a very high value for liquids (due to partial ionisation), and that hydrogen has a high dimensionless Prandtl group $C_p\mu/k$, which is approximately constant (where C_p is the specific heat at constant pressure and μ is the viscosity), can be used to evaluate k at high temperatures where it is difficult to determine a value experimentally because of the formation of convection currents. k does not vary significantly with pressure, except metallic solids lower values, non-metallic liquids low values, and gases very low values. It is important to note that amongst metals, stainless steel has a low value, that water value for gases (due to the high mobility of the molecules). With gases, k decreases with where this is reduced to a value so low that the mean free path of the molecules becomes

HEAT TRANSFER

389

4	15	Дешь		K	Υ	Temp	
(W/mK)	(Btu/h ft ² °F/ft)	(K)	N	(M/mK)	(Biu/h fi ² °F/fi)	(K)	
S£.0	02.0	563	sbiupi.d 2002 bigg 2iftaA	UEC.	133	CLS	Solids — Metals
71.0	01.0	£0£	Acetic acid 50%	730	133	ELS	muinimulA
. 71.0			Acetone	#6	7 S	167	Cadmium
91.0	1.0 60.0	273–293	AnilinA	LLE	218	ELE	Copper
01.0	60:0	303	Benzene Calcium chloride	87 19	85.72	97E 16Z	Iron (wrought)
cc. 0	25.0	303	%0£ annd	04	0:17	076	Iron (cast)
0.24	7£1.0	793	Ethyl alcohol 80%	. EE	16	ETE	hea I
8£.0	22.0	793	Glycerol 60%	LS	33	575	Lead Nickel
24.0	92.0	763	Glycerol 40%	415	238	£7£	
41.0	80.0	303	n-Heptane	St	97 967	767	Silver Sieel 1% C
9£.8	4.83	301	Mercury	55	35	167	
96.0	12.0	303	Sulphuric acid 90%	cc	70	1.67	Tantalum
64.0	22.0	303	%00 bios arinquis	113	\$9	303	fetam vtferimhA
79'0	955.0	303	Water	681	601		Admiralty metal
99.0	186.0	EEE .	Water	91	. 2.6	263	Bronze Stainless Steel
			Gases				solids — Non-metals
71.0	01.0	273	Hydrogen	71.0	960.0	373	Asbestos sheet
\$10.0	2800.0	273	Carbon dioxide	91.0	60.0	273	Asbestos
420.0	0.014	£L7	тiА	61.0	11.0	£ L £	Asbestos
1.50.0	810.0	ELE	τiA	12.0	0.12	€74	Aspestos
670.0	710.0	273	Methane	1.£	8.1	£01	Bricks (alumina)
0.025	0.0145	ELE	Water vapour	69.0	4.0	. 293	Bricks (building)
₽20.0	8£10.0	273	Мітовеп	8.£	2.2	£L\$	Magnesite
710.0	L600'0	273	Ethylene	0.050	670.0	303	Cotton wool
0.024	1410.0	273	Oxygen	1.09	£9.0	303	Glass
810.0	9010.0	273	Ethane	64.0	22.0	373	Mica
				61.0	780.0	£72 .	Кпррст (рагд)
		•		280.0	€0.0	263	Sawdust
				640.0	220.0	303	Cork
				140.0	0.024		Glass wool
				0.070	≯ 0.0	_	85% Magnesia
				121	L8	273	Graphite

391

HEAT TRANSFER

comparable with the dimensions of the vessel; further reduction of pressure then causes k to decrease.

Typical values for Prandtl numbers are as follows:

_		Glycerol 1000	nelts 1	
٠		1.38 Glyc		Men
Air	Oxygen	Ammonia (gas)	Water	

The low conductivity of heat insulating materials, such as cork, glass wool, and so on, is largely accounted for by their high proportion of air space. The flow of heat through such materials is governed mainly by the resistance of the air spaces, which should be sufficiently small for convection currents to be suppressed.

It is convenient to rearrange equation 9.12 to give:

$$Q = \frac{(T_1 - T_2)A}{(x/k)} \tag{9.17}$$

where x/k is known as the thermal resistance and k/x is the transfer coefficient.

Example 9.2

Estimate the heat loss per square metre of surface through a brick wall 0.5 m thick when the inner surface is at 400 K and the outside surface is at 300 K. The thermal conductivity of the brick may be taken as 0.7 W/mK.

Solution

From equation 9.12:

$$Q = \frac{0.7 \times 1 \times (400 - 300)}{0.5} \qquad \frac{W}{\text{PM/K}} = \frac{L_{\text{M}}}{M^2}$$

9.3.2. Thermal resistances in series

It has been noted earlier that thermal resistances may be added together for the case of heat transfer through a complete section formed from different media in series.

Figure 9.6 shows a composite wall made up of three materials with thermal conductivities k_1 , k_2 , and k_3 , with thicknesses as shown and with the temperatures T_1 , T_2 , T_3 , and T_4 at the faces. Applying equation 9.12 to each section in turn, and noting that the same quantity of heat Q must pass through each area A:

$$T_1 - T_2 = \frac{x_1}{k_1 A} Q, \ T_2 - T_3 = \frac{x_2}{k_2 A} Q \text{ and } T_3 - T_4 = \frac{x_3}{k_3 A} Q$$
On addition:
$$(T_1 - T_4) = \left(\frac{x_1}{k_1 A} + \frac{x_2}{k_2 A} + \frac{x_3}{k_3 A} \right) Q$$
 (9.18)

Figure 9.6. Conduction of heat through a composite wall

$$Q = \frac{T_1 - T_4}{\Sigma(x_1/k_1A)}$$
= Total driving force
$$= \frac{\text{Total (thermal resistance/area)}}{\text{Total (thermal resistance/area)}}$$

or:

(9.19)

Example 9.3

A furnace is constructed with 0.20 m of firebrick, 0.10 m of insulating brick, and 0.20 m of building brick. The inside temperature is 1200 K and the outside temperature is 330 K. If the thermal conductivities are as shown in Figure 9.7, estimate the heat loss per unit area and the temperature at the junction of the firebrick and the insulating brick.

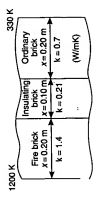


Figure 9.7. Data for Example 9.3

Solution

From equation 9.19:

$$Q = (1200 - 330) / \left[\left(\frac{0.20}{1.4 \times 1} \right) + \left(\frac{0.10}{0.21 \times 1} \right) + \left(\frac{0.20}{0.7 \times 1} \right) \right]$$

$$= \frac{870}{(0.143 + 0.476 + 0.286)} = \frac{870}{0.905}$$

$$= \frac{961 \text{ W/m}^2}{1.000}$$

The ratio (Temperature drop over firebrick)/(Total temperature drop) = (0.143/0.905)

Temperature drop over firebrick =
$$\left(\frac{870 \times 0.143}{0.905}\right) = 137 \text{ deg K}$$

Hence the temperature at the firebrick-insulating brick interface = (1200 - 137) = 1063 K

9.3.3. Conduction through a thick-walled tube

The conditions for heat flow through a thick-walled tube when the temperatures on the inside and outside are held constant are shown in Figure 9.8. Here the area for heat flow is proportional to the radius and hence the temperature gradient is inversely proportional

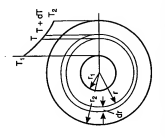


Figure 9.8. Conduction through thick-walled tube or spherical shell

The heat flow at any radius r is given by:

$$Q = -k2\pi r l \frac{dT}{dr} \tag{9.20}$$

where l is the length of tube.

Integrating between the limits r₁ and r₂:

$$Q \int_{r_1}^{r_2} \frac{\mathrm{d}r}{r} = -2\pi lk \int_{r_1}^{r_2} \mathrm{d}T$$

$$Q = \frac{2\pi l k(T_1 - T_2)}{\ln(r_1/r_1)} \tag{9.21}$$

This equation may be put into the form of equation 9.12 to give:

ö

$$Q = \frac{k(2\pi r_m l)(T_1 - T_2)}{r_2 - r_1}$$
(9.22)

where $r_m = (r_2 - r_1)/\ln(r_2/r_1)$, is known as the *logarithmic mean radius*. For thin-walled tubes the arithmetic mean radius r_a may be used, giving:

$$=\frac{k(2\pi r_a l)(T_1 - T_2)}{r_2 - r_1}$$
(9.23)

9.3.4. Conduction through a spherical shell and to a particle

For heat conduction through a spherical shell, the heat flow at radius r is given by:

$$Q = -k4\pi r^2 \frac{dT}{dr} \tag{9.24}$$

HEAT TRANSFER

Integrating between the limits r₁ and r₂:

$$Q \int_{r_1}^{r_2} \frac{dr}{r^2} = -4\pi k \int_{r_1}^{r_2} dT$$

$$Q = \frac{4\pi k (T_1 - T_2)}{(1/r_1) - (1/r_2)}$$
(9.

An important application of heat transfer to a sphere is that of conduction through a example in fluidised beds, rotary kilns, spray dryers and plasma devices. If the temperature stationary fluid surrounding a spherical particle or droplet of radius r as encountered for difference $T_1 - T_2$ is spread over a very large distance so that $r_2 = \infty$ and T_1 is the temperature of the surface of the drop, then:

$$\frac{Qr}{(4\pi r^2)(T_1 - T_2)k} = 1$$

$$\frac{hd}{k} = Nu' = 2$$
(9.26)

ë.

where $Q/4\pi r^2(T_1-T_2)=h$ is the heat transfer coefficient, d is the diameter of the particle or droplet and hd/k is a dimensionless group known as the Nusselt number (Nu')for the particle. The more general use of the Nusselt number, with particular reference to neat transfer by convection, is discussed in Section 9.4. This value of 2 for the Nusselt number is the theoretical minimum for heat transfer through a continuous medium. It is greater if the temperature difference is applied over a finite distance, when equation 9.25 must be used. When there is relative motion between the particle and the fluid the heat transfer rate will be further increased, as discussed in Section 9.4.6.

from the surface to the internal parts of the particle is of importance. For example, if the particle is a poor conductor then the rate at which the particulate material reaches some desired average temperature may be limited by conduction inside the particle rather than In this approach, heat transfer to a spherical particle by conduction through the surrounding fluid has been the prime consideration. In many practical situations the flow of heat by conduction to the outside surface of the particle. This problem involves unsteady state transfer of heat which is considered in Section 9.3.5.

Equations may be developed to predict the rate of change of diameter d of evaporating stagnant gas to the droplet surface, and heat transfer is the rate controlling step, it is practical problem is the prediction of the residence time required in a combustion chamber back from the flame to the droplet surface provides the heat to vaporise the oil and sustain the surrounding flame. Again d^2 decreases approximately linearly with time though the droplets. If the latent heat of vaporisation is provided by heat conducted through a hotter shown by SPALDING⁽²⁾ that d² decreases linearly with time. A closely related and important to ensure virtually complete burning of the oil droplets. Complete combustion is desirable to obtain maximum utilisation of energy and to minimise pollution of the atmosphere by derivation of the equation is more complex due to mass transfer effects, steep temperature partially burned oil droplets. Here a droplet is surrounded by a flame and heat conducted gradients⁽³⁾ and circulation in the drop⁽⁴⁾.

9.3.5. Unsteady state conduction

Basic considerations

In the problems which have been considered so far, it has been assumed that the conditions at any point in the system remain constant with respect to time. The case of heat transfer by conduction in a medium in which the temperature is changing with time is now considered. This problem is of importance in the calculation of the temperature distribution in a body which is being heated or cooled. If, in an element of dimensions dx by dy by dz (Figure 9.9), the temperature at the point (x, y, z) is θ and at the point (x + dx, y + dy,z + dz) is $(\theta + d\theta)$, then assuming that the thermal conductivity k is constant and that no heat is generated in the medium, the rate of conduction of heat through the element is:

$$= -k \, dy \, dz \left(\frac{\partial \theta}{\partial x}\right)_{yz} \quad \text{in the } x\text{-direction}$$

$$= -k \, dz \, dx \left(\frac{\partial \theta}{\partial y}\right)_{zx} \quad \text{in the } y\text{-direction}$$

$$= -k \, dx \, dy \left(\frac{\partial \theta}{\partial z}\right)_{xy} \quad \text{in the } z\text{-direction}$$

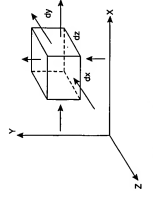


Figure 9.9. Element for heat conduction

The rate of change of heat content of the element is equal to *minus* the rate of increase of heat flow from (x, y, z) to (x + dx, y + dy, z + dz). Thus the rate of change of the heat content of the element is:

$$= k \, \mathrm{d}y \, \mathrm{d}z \left(\frac{\partial^2 \theta}{\partial x^2} \right)_{yz} \, \mathrm{d}x + k \, \mathrm{d}z \, \mathrm{d}x \left(\frac{\partial^2 \theta}{\partial y^2} \right)_{zx} \, \mathrm{d}y + k \, \mathrm{d}x \, \mathrm{d}y \left(\frac{\partial^2 \theta}{\partial z^2} \right)_{xy} \, \mathrm{d}z$$

$$= k \, \mathrm{d}x \, \mathrm{d}y \, \mathrm{d}z \left[\left(\frac{\partial^2 \theta}{\partial x^2} \right)_{yz} + \left(\frac{\partial^2 \theta}{\partial y^2} \right)_{zz} + \left(\frac{\partial^2 \theta}{\partial z^2} \right)_{zy} \right] \tag{9.27}$$

The rate of increase of heat content is also equal, however, to the product of the heat capacity of the element and the rate of rise of temperature.

and:

Thus:
$$k \, dx \, dy \, dz \left[\left(\frac{\partial^2 \theta}{\partial x^2} \right)_{yz} + \left(\frac{\partial^2 \theta}{\partial y^2} \right)_{zz} + \left(\frac{\partial^2 \theta}{\partial z^2} \right)_{xy} \right] = C_p \rho \, dx \, dy \, dz \, \frac{\partial \theta}{\partial t}$$
or:
$$\frac{\partial \theta}{\partial t} = \frac{k}{C_p \rho} \left[\left(\frac{\partial^2 \theta}{\partial x^2} \right)_{yz} + \left(\frac{\partial^2 \theta}{\partial y^2} \right)_{xz} + \left(\frac{\partial^2 \theta}{\partial z^2} \right)_{xy} \right]$$

$$= D_H \left[\left(\frac{\partial^2 \theta}{\partial x^2} \right)_{yz} + \left(\frac{\partial^2 \theta}{\partial y^2} \right)_{xz} + \left(\frac{\partial^2 \theta}{\partial z^2} \right)_{xy} \right]$$
(9.28)

where $D_H = k/C_p \rho$ is known as the *thermal diffusivity*. This partial differential equation is most conveniently solved by the use of the Laplace transform of temperature with respect to time. As an illustration of the method of solution, the problem of the unidirectional flow of heat in a continuous medium will be considered The basic differential equation for the X-direction is:

$$\frac{\partial \theta}{\partial t} = D_H \frac{\partial^2 \theta}{\partial x^2} \tag{9.29}$$

This equation cannot be integrated directly since the temperature θ is expressed as a involves transforming the equation so that the Laplace transform of θ with respect to time is used in place of θ . The equation then involves only the Laplace transform $\bar{\theta}$ and the distance x. The Laplace transform of θ is defined by the relation: function of two independent variables, distance x and time t. The method of solution

$$\overline{\theta} = \int_0^\infty \theta e^{-\mu t} dt \tag{9.3}$$

where p is a parameter. Thus $\bar{\theta}$ is obtained by operating on θ with respect to t with x constant.

Then:
$$\frac{\partial^2 \overline{\theta}}{\partial x^2} = \frac{\partial^2 \overline{\theta}}{\partial x^2}$$
 and:
$$\frac{\partial \overline{\theta}}{\partial t} = \int_0^\infty \frac{\partial \theta}{\partial t} e^{-\rho t} dt$$
$$= \left[\theta e^{-\rho t} \right]_0^\infty + \rho \int_0^\infty e^{-\rho t} \theta dt$$
$$= -\theta_{t=0} + \rho \overline{\theta}$$
 (9.32)

Then, taking the Laplace transforms of each side of equation 9.29:

$$\frac{\partial \theta}{\partial t} = D_H \frac{\partial^2 \theta}{\partial x^2}$$

$$p\bar{\theta} - \theta_{t=0} = D_H \frac{\partial^2 \bar{\theta}}{\partial x^2} \quad \text{(from equations 9.31 and 9.32)}$$

$$\frac{\partial^2 \bar{\theta}}{\partial x^2} - \frac{p}{D_H} \bar{\theta} = -\frac{\theta_{t=0}}{D_H}$$

or:

HEAT TRANSFER

$$\bar{\theta} = B_1 e^{\sqrt{(p/D_H)^3}} + B_2 e^{-\sqrt{(p/D_H)^3}} + \theta_{i=0} p^{-1}$$
 (9.3)

and therefore:
$$\frac{\partial \overline{\theta}}{\partial x} = B_1 \sqrt{\frac{\overline{p}}{D_H}} e^{\sqrt{(\overline{p}/\overline{D}_H)}x} - B_2 \sqrt{\frac{\overline{p}}{D_H}} e^{-\sqrt{(\overline{p}/\overline{D}_H)}x}$$
 (9.34)

The temperature θ , corresponding to the transform $\bar{\theta}$, may now be found by reference to tables of the Laplace transform. It is first necessary, however, to evaluate the constants B₁ and B₂ using the boundary conditions for the particular problem since these constants will in general involve the parameter p which was introduced in the transformation.

Considering the particular problem of the unidirectional flow of heat through a body with plane parallel faces a distance l apart, the heat flow is normal to these faces and the temperature of the body is initially constant throughout. The temperature scale will be so chosen that this uniform initial temperature is zero. At time, t = 0, one face (at x = 0) will be brought into contact with a source at a constant temperature θ' and the other face (at x = l) will be assumed to be perfectly insulated thermally

The boundary conditions are therefore:

$$t = 0, \qquad \theta = 0$$

$$t = 0$$
, $\theta = 0$
 $t > 0$, $\theta = \theta$ when $x = 0$

$$t > 0$$
, $\frac{\partial \theta}{\partial x} = 0$ when $x = l$

$$t > 0$$
, $\frac{1}{\partial x} = 0$ when $x = \frac{1}{6}$

$$\overline{\theta}_{x=0} = \int_0^\infty \theta' e^{-pt} dt = \frac{\theta'}{p}$$

$$(3\overline{\theta})$$

Substitution of these boundary conditions in equations 9.33 and 9.34 gives:

$$B_1 + B_2 = \frac{\theta'}{p}$$

$$B_1 e^{\sqrt{(\rho/D_H)}l} - B_2 e^{-\sqrt{(\rho/D_H)}l} = 0$$
 (9.35)

Hence:

$$B_1 = \frac{(\theta/p)e^{-\sqrt{(p/D_H)^2}}}{e^{\sqrt{(p/D_H)^2}} + e^{-\sqrt{(p/D_H)^2}}}$$

$$B_2 = \frac{(\theta'/p)e^{\sqrt{(p/D_H)l}}}{e^{\sqrt{(p/D_H)l}} + e^{-\sqrt{(p/D_H)l}}}$$

 $= \frac{\theta'}{p} (e^{(l-x)} \sqrt{(p/DH)} + e^{-(l-x)} \sqrt{(p/DH)}) (1 + e^{-2} \sqrt{(p/DH)^l})^{-1} (e^{-\sqrt{(p/DH)^l}})$ $\frac{e^{(l-x)\sqrt{(p/D_H)}} + e^{-(l-x)\sqrt{(p/D_H)}} \theta}{e^{\sqrt{(p/D_H)^l}} + e^{-\sqrt{(p/D_H)^l}}} \frac{\theta}{P}$

 $= \frac{\theta'}{p} (e^{-x} \sqrt{(p/D_H)} + e^{-(2l-x)} \sqrt{(p/D_H)}) (1 - e^{-2l} \sqrt{(p/D_H)} + \cdots + (-1)^N e^{-2Nl} \sqrt{(p/D_H)} + \cdots)$

$$= \sum_{N=0}^{N=\infty} \frac{\theta'}{P} (-1)^N (e^{-(2IN+x)\sqrt{(P/DH)}} + e^{-(2(N+1)I-x)\sqrt{(P/DH)}})$$
 (9.36)

The temperature θ is then obtained from the tables of inverse Laplace transforms in the Appendix (Table 12, No 83) and is given by:

$$\theta = \sum_{N=0}^{N=\infty} (-1)^N \theta' \left(\text{erfc} \frac{2lN + x}{2\sqrt{D_H t}} + \text{erfc} \frac{2(N+1)l - x}{2\sqrt{D_H t}} \right)$$
 (9.3)

$$\operatorname{erfc} x = \frac{2}{\sqrt{\pi}} \int_{x}^{\infty} e^{-\xi^{2}} d\xi$$

 $\sqrt{\pi} \ J_x$ Values of erfc x (= 1 – erf x) are given in the Appendix (Table 13) and in specialist sources.⁽⁵⁾

Equation 9.37 may be written in the form:
$$\frac{\theta}{\theta} = \sum_{N=0}^{N=\infty} (-1)^N \left\{ \text{erfc} \left[Fo_l^{-1/2} \left(N + \frac{1}{2} \frac{x}{l} \right) \right] + \text{erfc} \left[Fo_l^{-1/2} \left((N+1) - \frac{1}{2} \frac{x}{l} \right) \right] \right\}$$

where $Fo_l = (D_H t/l^2)$ and is known as the Fourier number.

$$\frac{\theta}{\theta} = f\left(Fo_l, \frac{x}{l}\right) \tag{9.39}$$

The numerical solution to this problem is then obtained by inserting the appropriate values for the physical properties of the system and using as many terms in the series as are necessary for the degree of accuracy required. In most cases, the above series converge quite rapidly.

is applicable when there are discontinuities in the physical properties of the material. (6) The boundary conditions, however, become a little more complicated, but the problem is This method of solution of problems of unsteady flow is particularly useful because it intrinsically no more difficult.

consists of replacing the heat flow problem by the analogous electrical situation and measuring the electrical potentials at various points. The heat capacity per unit volume $C_p \rho$ is represented by an electrical capacitance, and the thermal conductivity k by an A general method of estimating the temperature distribution in a body of any shape

electrical conductivity. This method can be used to take account of variations in the thermal properties over the body.

Example 9.4

Calculate the time taken for the distant face of a brick wall, of thermal diffusivity $D_H = 0.0043 \text{ cm}^2/s$ and thickness l = 0.45 m, to rise from 295 to 375 K, if the whole wall is initially at a constant temperature of 295 K and the near face is suddenly raised to 900 K and maintained at this temperature. Assume that all the flow of heat is perpendicular to the faces of the wall and that the distant face is perfectly insulated.

Solution

The temperature at any distance x from the near face at time t is given by:

$$\theta = \sum_{N=0}^{N=\infty} (-1)^N \theta' \left\{ \text{erfc} \left[\frac{2!N+x}{2\sqrt{D_H t}} \right] + \text{erfc} \left[\frac{2(N+1)!-x}{2\sqrt{D_H t}} \right] \right\}$$
 (equation 9.37)

The temperature at the distant face is therefore given by

$$\theta = \sum_{N=0}^{N=\infty} (-1)^N \theta' 2 \text{ erfc} \left[\frac{(2N+1)l}{2\sqrt{D_H t}} \right]$$

Choosing the temperature scale so that the initial temperature is everywhere zero, then:

$$\frac{\theta}{2\theta} = \frac{375 - 295}{2(900 - 295)} = 0.066$$

$$D_H = 4.2 \times 10^{-7} \text{ m}^2 / s \quad .. \quad \sqrt{D_H} = 6.5 \times 10^{-4}$$

$$I = 0.45 \text{ m}$$

$$I = 0.45 \text{$$

$$= \sum_{N=0}^{N=\infty} (-1)^N \operatorname{crfc} \left[\frac{346(2N+1)}{t^{0.5}} \right]$$

= $erfc(346t^{-0.5}) - erfc(1038t^{-0.5}) + erfc(1730t^{-0.5}) - \cdots$

An approximate solution is obtained by taking the first term only, to give:

$$346t^{-0.5} = 1.30$$
$$t = 70\,840 \text{ s}$$

from which

= 70.8 ks or 19.7 h

Schmidt's method

Numerical methods have been developed by replacing the differential equation by a finite difference equation. Thus in a problem of unidirectional flow of heat:

$$\frac{\partial \theta}{\partial t} \approx \frac{\theta_{x(t+\Delta t)} - \theta_{x(t-\Delta t)}}{2\Delta t} \approx \frac{\theta_{x(t+\Delta t)} - \theta_{xt}}{\Delta t}$$

HEAT TRANSFER

$$\frac{\partial^2 \theta}{\partial x^2} \approx \frac{\left(\frac{\theta_{(x+\Delta x)t} - \theta_{xt}}{\Delta x} - \frac{\theta_{xt} - \theta_{(x-\Delta x)t}}{\Delta x}\right)}{\Delta x}$$

$$= \frac{\theta_{(x+\Delta x)t} + \theta_{(x-\Delta x)t} - 2\theta_{xt}}{(\Delta x)^2}$$

where θ_{x} is the value of θ at time t and distance x from the surface, and the other values of θ are at intervals Δx and Δt as shown in Figure 9.10.

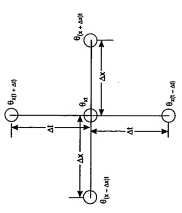


Figure 9.10. Variation of temperature with time and distance

Substituting these values in equation 9.29:

$$\theta_{x(t+\Delta t)} - \theta_{x(t-\Delta t)} = D_H \frac{2\Delta t}{(\Delta x)^2} (\theta_{(x+\Delta x)t} + \theta_{(x-\Delta x)t} - 2\theta_H)$$
 (9.40)

$$\theta_{x(t+\Delta t)} - \theta_{xt} = D_H \frac{\Delta t}{(\Delta x)^2} (\theta_{(x+\Delta x)t} + \theta_{(x-\Delta x)t} - 2\theta_{xt})$$
 (9.41)

and:

at time $t + \Delta t$ can be calculated by the application of equation 9.41 over the whole extent of the body in question. The intervals Δx and Δt are so chosen that the required degree Thus, if the temperature distribution at time t, is known, the corresponding distribution of accuracy is obtained.

A graphical method of procedure has been proposed by SCHMIDT⁽⁷⁾. If the temperature distribution at time t is represented by the curve shown in Figure 9.11 and the points representing the temperatures at $x - \Delta x$ and $x + \Delta x$ are joined by a straight line, then the distance θ_a is given by:

$$\theta_a = \frac{\theta_{(x+\Delta x)t} + \theta_{(x-\Delta x)t}}{2} - \theta_{xt}$$

$$= \frac{(\Delta x)^2}{2D_H \Delta t} (\theta_{x(t+\Delta t)} - \theta_{xt}) \cdot (\text{from equation 9.41})$$
 (9.4)

900 K

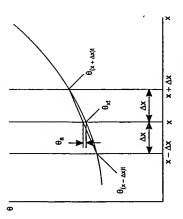


Figure 9.11. Schmidt's method

Thus, θ_a represents the change in θ_{xt} after a time interval Δt , such that:

$$\Delta t = \frac{(\Delta x)^2}{2D_H} \tag{9.43}$$

295--냒

375--

If this simple construction is carried out over the whole of the body, the temperature distribution after time Δt is obtained. The temperature distribution after an interval $2\Delta t$ is then obtained by repeating this procedure.

The most general method of tackling the problem is the use of the *finite-element* technique⁽⁸⁾ to determine the temperature distribution at any time by using the finite difference equation in the form of equation 9.40.

Example 9.5

Solve Example 9.4 using Schmidt's method.

Solution

The development of the temperature profile is shown in Figure 9.12. At time t = 0 the temperature is constant at 295 K throughout and the temperature of the hot face is raised to 900 K. The problem will be solved by taking relatively large intervals for Δx .

Choosing $\Delta x = 50$ mm, the construction shown in Figure 9.12 is carried out starting at the bot face. Points corresponding to temperature after a time interval Δt are marked 1, after a time interval $2\Delta t$ by 2,

Points corresponding to temperature after a time interval Δt are marked 1, after a time interval $2\Delta t$ by 2, and so on. Because the second face is perfectly insulated, the temperature gradient must be zero at this point. Thus, in obtaining temperatures at x = 450 mm it is assumed that the temperature at x = 500 mm will be the same as at x = 400 mm, that is, horizontal lines are drawn on the diagram. It is seen that the temperature is less than 375 K after time $23\Delta t$ and greater than 375 K after time $25\Delta t$.

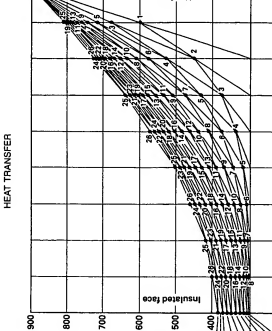
From equation 9.43:
$$\Delta t = 5.0^2/(2 \times 0.0042) = 2976 \text{ s}$$

Thus time required
$$= 24 \times 2976 = 71400 \text{ s}$$

$$71.4 \text{ ks} = 19.8 \text{ h}$$

ë

This value is quite close to that obtained by calculation, even using the coarse increments in Δx .



Hot face

Temperature (K)

Figure 9.12. Development of temperature profile

Distance from hot face (mm)

200

250

300

350

충 8

Heating and cooling of solids and particles

The exact mathematical solution of problems involving unsteady thermal conduction may be very difficult, and sometimes impossible, especially where bodies of irregular shapes are concerned, and other methods are therefore required.

When a body of characteristic linear dimension L, initially at a uniform temperature θ_0 , is exposed suddenly to surroundings at a temperature θ' , the temperature distribution at any time t is found from dimensional analysis to be:

$$\frac{\theta' - \theta}{\theta' - \theta_0} = f\left(\frac{hL}{k}, D_H \frac{t}{L^2}, \frac{x}{L}\right) \tag{9.44}$$

where D_H is the thermal diffusivity $(k_p/C_p\rho)$ of the solid, x is distance within the solid body and h is the heat transfer coefficient in the fluid at the surface of the body.

Analytical solutions of equation 9.44 in the form of infinite series are available for some simple regular shapes of particles, such as rectangular slabs, long cylinders and spheres, for conditions where there is heat transfer by conduction or convection to or from the surrounding fluid. These solutions tend to be quite complex, even for simple shapes. The heat transfer process may be characterised by the value of the *Biot number Bi* where:

$$Bi = \frac{hL}{k_p} = \frac{L/k_p}{1/h} \tag{9.45}$$

L is a characteristic dimension, such as radius in the case of a sphere or long cylinder, or half the thickness in the case of a slab, and

 k_p is the thermal conductivity of the particle.

The Biot number is essentially the ratio of the resistance to heat transfer within the particle to that within the external fluid. At first sight, it appears to be similar in form to the Nusselt Number Nu' where:

$$Nu' = \frac{hd}{k} = \frac{2hr_o}{k} \tag{9.46}$$

However, the Nusselt number refers to a single fluid phase, whereas the Biot number is related to the properties of both the fluid and the solid phases.

Three cases are now considered:

- (1) Very large Biot numbers, Bi → ∞
 (2) Very low Biot numbers, Bi → 0
 (3) Intermediate values of the Biot number.

(1) Bi very large. The resistance to heat transfer in the fluid is then low compared with equal to the bulk temperature of the fluid, and the heat transfer rate is independent of the that in the solid with the temperature of the surface of the particle being approximately Biot number. Equation 9.44 then simplifies to:

$$\frac{\theta' - \theta}{\theta' - \theta_0} = f\left(D_H \frac{t}{L^2}, \frac{x}{L}\right) = f\left(Fo_L, \frac{x}{L}\right) \tag{9.47}$$

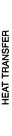
where $Fo_L\left(=D_H\frac{t}{L^2}\right)$ is known as the Fourier number, using L in this case to denote the characteristic length, and x is distance from the centre of the particle. Curves connecting these groups have been plotted by a number of workers for bodies of various shapes, although the method is limited to those shapes which have been studied experimentally.

In Figure 9.13, taken from CARSLAW and JAEGER⁽⁵⁾, the value of $(\theta' - \theta_c)/(\theta' - \theta_0)$ is plotted to give the temperature θ_c at the centre of bodies of various shapes, initially at a uniform temperature θ_0 , at a time t after the surfaces have been suddenly altered to and maintained at a constant temperature θ' .

In this case (x/L) is constant at 0 and the results are shown as a function of the particular value of the Fourier number Fo_L $(D_H t/L^2)$.

fluid; this occurs when the thermal conductivity of the particle in very high and/or when uniform and a "lumped capacity" analysis may be performed. Thus, if a solid body of (2) Bi very small. (say, <0.1). Here the main resistance to heat transfer lies within the the particle is very small. Under these conditions, the temperature within the particle is volume V and initial temperature θ_0 is suddenly immersed in a volume of fluid large enough for its temperature θ to remain effectively constant, the rate of heat transfer from the body may be expressed as:

$$-\rho C_p V \frac{d\theta}{dt} = h A_e (\theta - \theta') \tag{9.48}$$



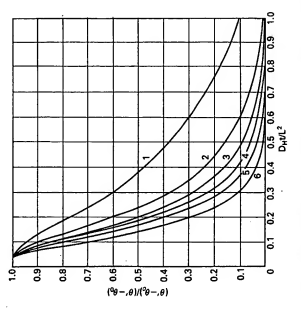


Figure 9.13. Cooling curve for bodies of various shapes: 1, slab (2L = thickness); 2, square bar (2L = side); 3, long cylinder (L = radius); 4, cube (2L = length of side); 5, cylinder (L = radius, length = 2L); 6, sphere (L = radius)

where A_e is the external surface area of the solid body.

Then:
$$\int_{\theta_0}^{\theta} \frac{d\theta}{\theta - \theta'} = -\int_0^t \frac{hA_e}{\rho C_v V} dt$$
i.e.:
$$\frac{\theta - \theta'}{\theta_0 - \theta'} = e^{-t/\tau}$$

(9.49)

where $\tau = \frac{\rho C_P V}{h A_e}$ is known as the *response time constant*. It will be noted that the relevant characteristic dimension in the Biot number is defined as the ratio of the volume to the external surface area of the particle (V/A_e) , and the higher the value of V/A_e , then the slower will be the response time. With the characteristic dimension defined in this way, this analysis is valid for particles of any shape at values of the Biot number less than 0.1

Example 9.6

A 25 mm diameter copper sphere and a 25 mm copper cube are both heated in a furnace to 650 °C (923 K). They are then annealed in air at 95 °C (368 K). If the external heat transfer coefficient h is 75 W/m²K in both cases, what is temperature of the sphere and of the cube at the end of 5 minutes?

4

The physical properties at the mean temperature for copper are:

 $\rho = 8950 \text{ kg/m}^3$ $C_p = 0.38 \text{ kJ/kg K}$ $k_p = 385 \text{ W/mK}$

Solution

$$V/A_e$$
 for the sphere
$$= \frac{\frac{\pi}{6}d^3}{\pi d^2} = \frac{d}{6} = \frac{25 \times 10^{-3}}{6} = 4.17 \times 10^{-3} \text{ m}$$

$$V/A_e \text{ for the cube} = \frac{l^3}{6l^2} = \frac{l}{6} = \frac{25 \times 10^{-3}}{6} = 4.17 \times 10^{-3} \text{ m}$$

$$Bi = \frac{h(V/A_e)}{k} = \frac{75 \times 25 \times 10^{-3}}{385 \times 6} = 8.1 \times 10^{-4} \ll 0.1$$

The use of a lumped capacity method is therefore justified.

$$\tau = \frac{\rho C_p V}{h A_e} = \frac{8950 \times 380}{75} \times \frac{25 \times 10^{-3}}{6} = 189 \text{ s}$$

Then using equation 9.49:

$$\frac{\theta - 368}{923 - 368} = \exp\left(-\frac{5 \times 60}{189}\right)$$

 $\theta = 368 + 0.2045(923 - 368) = 481 \text{ K} = 208^{\circ}\text{C}$

Since the sphere and the cube have the same value of V/A_e , after 5 minutes they will both attain a temperature

(3) Intermediate values of Bi. In this case the resistances to heat transfer within the solid solid be uniform (case 1), nor will the surface temperature be equal to that in the bulk of body and the fluid are of comparable magnitude. Neither will the temperature within the

Analytical solutions in the form of infinite series can be obtained for some regular shapes (thin plates, spheres and long cylinders (length >> radius)), and numerical solutions and irregular. Some of the results have been presented by HEISLER⁽⁹⁾ in the form of charts, examples of which are shown in Figures 9.14-9.16 for thin slabs, long cylinders and spheres, respectively. It may be noted that in this case the characteristic length L is using finite element methods⁽⁸⁾ have been obtained for bodies of other shapes, both regular the half-thickness of the slab and the external radius r_o of the cylinder and sphere. the fluid (case 2)

centre-line or centre-point) to be obtained as a function of the Fourier number, and hence Figures 9.14–9.16 enable the temperature θ_c at the centre of the solid (centre-plane, of time, with the reciprocal of the Biot number (Bi^{-1}) as parameter.

Temperatures at off-centre locations within the solid body can then be obtained from a ture to the centre-temperature as a function of Biot number, with location within the particle or sphere). Additional charts are given by Heisler for the quantity of heat transferred from further series of charts given by Heisler (Figures 9.17-9.19) which link the desired temperaas parameter (that is the distance x from the centre plane in the slab or radius in the cylinder the particle in a given time in terms of the initial heat content of the particle.

Figures 9.17-9.19 clearly show that, as the Biot number approaches zero, the temperature becomes uniform within the solid, and the lumped capacity method may be used for calculating the unsteady-state heating of the particles, as discussed in section (2). The harts are applicable for Fourier numbers greater than about 0.2.

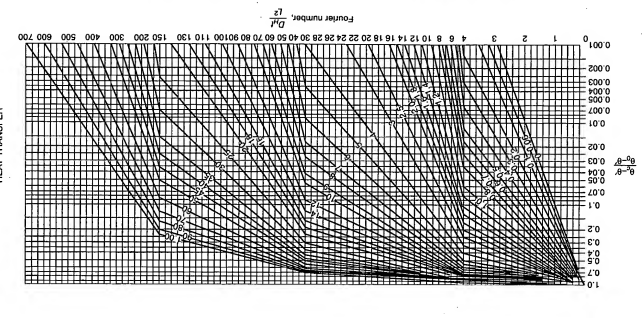


Figure 9.14. Mid-plane temperature for an infinite plate of thickness 2L, for various values of parameters $k_p/\hbar L (=Bi^{-1})$



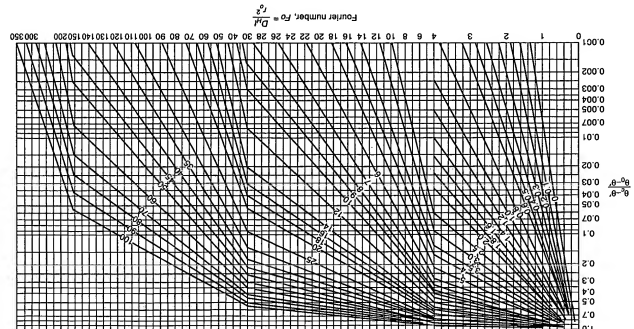


Figure 9.15. Axis temperature for an infinite cyclinder of radius v_o , for various of parameters $k_p/h v_o (= Bi^{-1})$

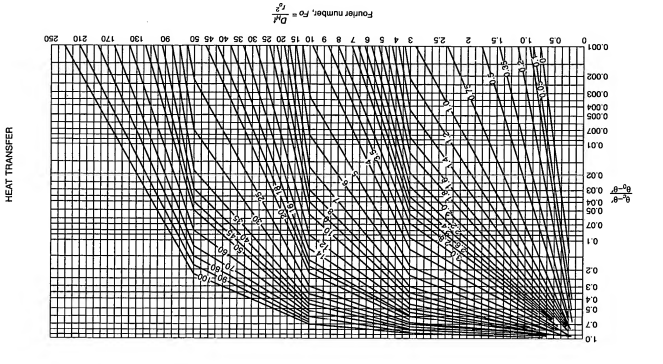


Figure 9.16. Centre-temperature for a sphere of radius r_o , for various values of parameters $\frac{k_p}{h r_o} (= Bi^{-1})$

HEAT TRANSFER

409

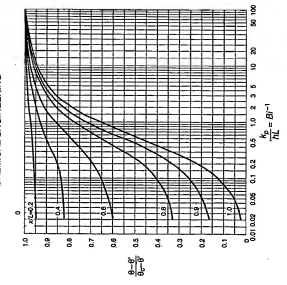


Figure 9.17. Temperature as a function of mid-plane temperature in an infinite plate of thickness 2L

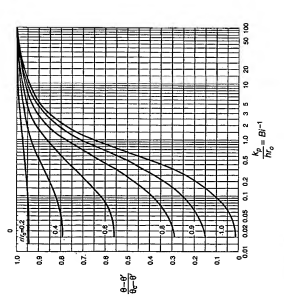


Figure 9.18. Temperature as a function of axis temperature in an infinite cylinder of radius r_o

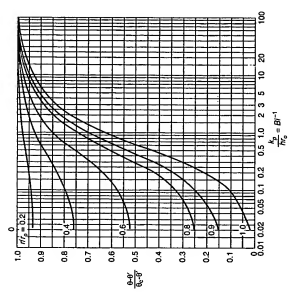


Figure 9.19. Temperature as a function of centre-temperature for a sphere of radius r_o

Example 9.7

A large thermoplastic sheet, 10 mm thick, at an initial temperature of 20° C (293 K), is to be heated in an oven in order to enable it to be moulded. The oven temperature is maintained at 100° C (373 K), the maximum temperature to which the plastic may be subjected, and it is necessary to ensure that the temperature throughout the whole of the sheet reaches a minimum of 80° C (353 K). Calculate the minimum length of time for which the sheet must be heated.

=2.5 W/mK Thermal conductivity kp of the plastic Thermal diffusivity of the surrounding fluid $D_H = 2 \times 10^{-7} \text{ m}^2/\text{s}$

External heat transfer coefficient h

 $=100 \text{ W/m}^2\text{K}$

Solution

Throughout the heating process, the temperature within the sheet will be a minimum at the centre-plane (x = 0) and therefore the required time is that for the centre to reach 80 °C (353 K).

For this process, the Biot number
$$Bi = \frac{hL}{k_p} = \frac{100 \times 5 \times 10^{-3}}{2.5} = 0.2$$
 and $Bi^{-1} = 5$

(since L, the half-thickness of the plate is 5 mm)

The limiting value of
$$\frac{\theta' - \theta_c}{\theta' - \theta_0} = \frac{373 - 353}{373 - 293} = 0.25$$

411

From Figure 9.17, the Fourier number $\frac{D_{H}t}{L^2} \approx 7.7$

 $t = \frac{7.7 \times (5 \times 10^{-3})^2}{2 \times 10^{-7}} = 960 \text{ s or } \frac{16 \text{ minutes}}{16 \times 10^{-7}}$

Thus:

Heating and melting of fine particles

There are many situations in which particles are heated or cooled by a surrounding gas and these may be classified according to the degree of movement of the particle as follows:

Although most beds of particles involve relatively large particle diameters, such as in pebble bed units used for the transfer of heat from flue gases to the incoming air for example, smaller particles, such as sand, are used in beds and, again, these are mainly used for heat recovery. One such application is the heating and cooling of buildings in hotter climes where the cool nocturnal air is used to cool a bed of particles which is then used to cool the incoming air during the heat of the day as it enters a building. In this way, an almost constant temperature may be achieved in a given enclosed environment in spite of the widely fluctuating ambient condition. A similar system has been used in less tropical areas where it is necessary to maintain a constant temperature in an environment in which heat is generated, such as a telephone exchange, for example. Such systems have the merit of very low capital and modest operating costs and, in most cases, the resistance to heat transfer by conduction within the solids is not dissimilar to the resistance in the gas film surrounding the particles.

ii) Partial movement of particles

The most obvious example of a process in which particles undergo only limited movement is the fluidised bed which is discussed in some detail in Volume 2. Applications here involve, not only heating and cooling, but also drying as in the case of grain dryers for example, and on occasions, chemical reaction as, for example, with fluidised-bed combustion. In such cases, conditions in the bed may, to all intents and purposes, be regarded as steady-state, with unsteady-state conduction taking place only in the entering 'process stream' which, by and large, is only a small proportion of the total bed mass in the bed.

iii) Falling particles

Particles fall by gravity through either static or moving gas streams in rotary dryers, for example, but they also fall through heating or cooling gases in specially designed columns. Examples here include the cooling of sand after it has been dried—again recovering heat in the process—salt cooling and also the spray drying of materials such as detergents which are sprayed as a concentrated solution of the material at the top of the tower and emerge as a dry powder. A similar situation occurs in fertiliser production where solid particles or granules are obtained from a spray of the molten material by counter-flow against a cooling gas stream. Convection to such materials is discussed in Section 9.4.6

One important problem involving unsteady state conduction of heat to particles is in the melting of powders in plasma spraying⁽¹⁰⁾ where Biot numbers can range from 0.005 to 5. In this case, there is initially a very high relative velocity between the fluid and the powder. The plasmas referred to here are partially ionised gases with temperatures of around 10,000 K formed by electric discharges such as arcs. There is an increasing industrial use

to non-continuum conditions discussed later, the value of Nu' = hd/k is therefore often resistant ceramic layers on centrifugal pump impellers and other equipment prone to cles, the particle is quickly accelerated to essentially the same velocity as the plasma jet(2) which can be melted before it strikes the surface. In the absence of complications due of the technique of plasma spraying in which powders are injected into a high-velocity plasma jet so that they are both melted and projected at velocities of several hundred metres per second onto a surface. The molten particles with diameters typically of the order 10-100 μm impinge to form an integral layer on the surface. Applications include the building up of worn shafts of pumps, for example, and the deposition of erosionerosion damage. When a powder particle first enters the plasma jet, the relative velocity may be hundreds of metres per second and heat transfer to the particle is enhanced by convection, as discussed in Section 9.4.6. Often, and more particularly for smaller partiand conduction becomes the main mechanism of heat transfer from plasma to particle. From a design point of view, neglecting the convective contribution will ease calculations and give a more conservative and safer estimate of the size of the largest particle taken as 2, as in equation 9.26.

One complication which arises in the application of this equation to powder heating in high temperature plasmas lies in the dependence of k, the thermal conductivity of the gas or plasma surrounding the particle, on temperature. For example, the temperature of the particle surface may be 1000 K, whilst that of the plasma away from the particle may be about 10,000 K or even higher. The thermal conductivity of argon increases by a factor of about 20 over this range of temperature and that of nitrogen gas passes through a pronounced peak at about 7100 K due to dissociation–recombination effects. Thus, the temperature at which the thermal conductivity k is evaluated will have a pronounced effect on the value of the external heat transfer coefficient. A mean value of k would seem appropriate where:

$$(k)_{\text{mean}} = \frac{1}{T_2 - T_1} \int_{T_1}^{T_2} k \, dT$$
 (equation 9.16)

Some workers have correlated experimental data in terms of k at the arithmetic mean temperature, and some at the temperature of the bulk plasma. Experimental validation of the true effective thermal conductivity is difficult because of the high temperatures, small particle sizes and variations in velocity and temperature in plasma jets.

In view of the high temperatures involved in plasma devices and the dependence of radiation heat transfer on T^4 , as discussed in Section 9.5, it is surprising at first sight that conduction is more significant than radiation in heating particles in plasma spraying. The explanation lies in the small values of d and relatively high values of k for the gas, both of which contribute to high values of h for any given value of Nu'. Also the emissivities of most gases are, as seen later in Section 9.5, rather low.

In situations where the surrounding fluid behaves as a non-continuum fluid, for example at very high temperatures and/or at low pressures, it is possible for Nu' to be less than 2. A gas begins to exhibit non-continuum behaviour when the mean free path between collisions of gas molecules or atoms with each other is greater than about 1/100 of the characteristic size of the surface considered. The molecules or atoms are then sufficiently far apart on average for the gas to begin to lose the character of a homogeneous or continuum fluid which is normally assumed in the majority of heat transfer or fluid

dynamics problems. For example, with a particle of diameter 25 μ m as encountered in, for example, oil-burner sprays, pulverised coal flames, and in plasma spraying in air at room temperature and atmospheric pressure, the mean free path of gas molecules is about 0.06 μ m and the air then behaves as a continuum fluid. If, however, the temperature were say 1800 K, as in a flame, then the mean free path would be about 0.33 μ m, which is greater than 1/100 of the particle diameter. Non-continuum effects, leading to values of Nu' lower than 2 would then be likely according to theory^(11,12). The exact value of Nu' depends on the surface accommodation coefficient. This is a difficult parameter to measure for the examples considered here, and hence experimental confirmation of the theory is difficult. At the still higher temperatures that exist in thermal plasma devices, non-continuum effects should be more pronounced and there is limited evidence that values of Nu' below 1 are obtained⁽¹⁰⁾. In general, non-continuum effects, leading in particular to values of Nu' less than 2, would be more likely at high temperatures, low pressures, and small particle sizes. Thus, there is an interest in these effects in the aerospace industry and small particle sizes. Thus, there is an interest in these effects in rocket engine

9.3.6. Conduction with internal heat source

If an electric current flows through a wire, the heat generated internally will result in a temperature distribution between the central axis and the surface of the wire. This type of problem will also arise in chemical or nuclear reactors where heat is generated internally. It is necessary to determine the temperature distribution in such a system and the maximum temperature which will occur.

If the temperature at the surface of the wire is T_o and the rate of heat generation per unit volume is Q_G , then considering unit length of a cylindrical element of radius r, the heat generated must be transmitted in an outward direction by conduction so that:

$$-k2\pi r \frac{dT}{dr} = \pi r^2 Q_G$$
Hence:
$$\frac{dT}{dr} = -\frac{Q_G r}{2k}$$
 (9.50)

 $T = -\frac{Q_G r^2}{A_L} + C$

Integrating

 $T = T_o$ when $r = r_o$ the radius of wire and hence:

$$T = T_o + Q_G \frac{r_o^2 - r^2}{4k}$$

$$T - T_o = \frac{Q_G r_o^2}{4k} \left(1 - \frac{r^2}{r_o^2} \right)$$
 (9.51)

ö

This gives a parabolic distribution of temperature and the maximum temperature will occur at the axis of the wire where $(T-T_o)=Q_Gr_o^2/4k$. The arithmetic mean temperature difference, $(T-T_o)_{av}=Q_Gr_o^2/8k$.

Since $Q_G\pi r_o^2$ is the rate of heat release per unit length of the wire then, putting T_1 as the temperature at the centre:

$$T_1 - T_o = \frac{\text{rate of heat release per unit length}}{4\pi k}$$
 (9.52)

Example 9.8

A fuel channel in a natural uranium reactor is 5 m long and has a heat release of 0.25 MW. If the thermal conductivity of the uranium is 33 W/mK, what is the temperature difference between the surface and the centre of the uranium element, assuming that the heat release is uniform along the rod?

Solution

Heat release rate =
$$0.25 \times 10^6$$
 W
$$= \frac{0.25 \times 10^6}{5} = 5 \times 10^4 \text{ W/m}$$

Thus, from equation 9.52:
$$T_1 - T_0 = \frac{5 \times 10^4}{4\pi \times 33}$$
 = 121 deg K

It should be noted that the temperature difference is independent of the diameter of the fuel rod for a cylindrical geometry, and that the heat released per unit volume has been considered as being uniform.

In practice the assumption of the uniform heat release per unit length of the rod is not valid since the neutron flux, and hence the heat generation rate varies along its length. In the simplest case where the neutron flux may be taken as zero at the ends of the fuel element, the heat flux may be represented by a sinusoidal function, and the conditions become as shown in Figure 9.20.

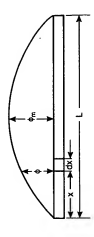


Figure 9.20. Variation of neutron flux along a length of fuel rod

Since the heat generated is proportional to the neutron flux, the heat dQ developed per unit time in a differential element of the fuel rod of length dx may be written as:

$$dQ = C \sin\left(\frac{\pi x}{t}\right) dx$$

The total heat generated by the rod Q is then given by:

$$Q = C \int_0^L \sin\left(\frac{\pi x}{L}\right) dx = \frac{2CL}{\pi}$$

Thus, $C = \pi Q/2L$. The heat release per unit length at any point is then given by:

$$\frac{\mathrm{d}Q}{\mathrm{d}x} = \frac{\pi Q}{2L} \sin\left(\frac{\pi x}{L}\right)$$

Substituting into equation 9.52 gives:

$$_{1}-T_{o}=\frac{\left(\frac{\pi Q}{2L}\right)\sin\left(\frac{\pi x}{L}\right)}{4\pi k}\tag{9.53}$$

It may be noted that when x = 0 or x = L, then $T_1 - T_0$ is zero as would be expected since the neutron flux was taken as zero at these positions.

9.4. HEAT TRANSFER BY CONVECTION

9.4.1. Natural and forced convection

than that of the remaining liquid. Cold liquid of higher density takes its place and a transfer process itself, natural convection occurs, such as in the heating of a vessel containing liquid by means of a heat source situated beneath it. The liquid at the bottom of the vessel becomes heated and expands and rises because its density has become less Heat transfer by convection occurs as a result of the movement of fluid on a macroscopic scale in the form of eddies or circulating currents. If the currents arise from the heat circulating current is thus set up.

In forced convection, circulating currents are produced by an external agency such as an agitator in a reaction vessel or as a result of turbulent flow in a pipe. In general, the magnitude of the circulation in forced convection is greater, and higher rates of heat transfer are obtained than in natural convection.

free of turbulence, covers the surface. In this film, heat transfer is by thermal conduction and, as the thermal conductivity of most fluids is low, the main resistance to transfer lies there. Thus an increase in the velocity of the fluid over the surface gives rise to the film coefficient increases as (fluid velocity)ⁿ, where 0.6 < n < 0.8, depending upon In most cases where convective heat transfer is taking place from a surface to a fluid, improved heat transfer mainly because the thickness of the film is reduced. As a guide, the circulating currents die out in the immediate vicinity of the surface and a film of fluid,

If the resistance to transfer is regarded as lying within the film covering the surface, the rate of heat transfer Q is given by equation 9.11 as:

$$Q = kA \frac{(T_1 - T_2)}{x}$$

The effective thickness x is not generally known and therefore the equation is usually rewritten in the form:

$$Q = hA(T_1 - T_2) (9.54)$$

where h is the heat transfer coefficient for the film and (1/h) is the thermal resistance.

HEAT TRANSFER

9.4.2. Application of dimensional analysis to convection

velocity of flow u of the fluid over the surface, the temperature difference ΔT and a factor determining the natural circulation effect caused by the expansion of the fluid on heating (the product of the coefficient of cubical expansion β and the acceleration due to individual effects by direct experimental methods. By arranging the variables in a series heat transfer rate per unit area q is dependent on those physical properties which affect flow pattern (viscosity μ and density ρ), the thermal properties of the fluid (the specific heat capacity C_p and the thermal conductivity k) a linear dimension of the surface l, the of dimensionless groups, however, the problem is made more manageable in that the number of groups is significantly less than the number of parameters. It is found that the So many factors influence the value of h that it is almost impossible to determine their gravity g). Writing this as a functional relationship:

$$q = \Phi[u, l, \rho, \mu, C_p, \Delta T, \beta g, k]$$

$$(9.55)$$

Noting the dimensions of the variables in terms of length L, mass M, time T, temperature θ , heat H:

$\mathrm{HL}^{-2}\mathrm{T}^{-1}$	LT^{-1}	Γ	$ML^{-1}T^{-1}$	ML-3	$\mathbf{H}\mathbf{L}^{-1}\mathbf{\Gamma}^{-1}\mathbf{\theta}^{-1}$	$HM^{-1}\theta^{-1}$	9		$\mathbf{L}\mathbf{L}^{-2}\theta^{-1}$
Heat transferred/unit area and unit time	Velocity	Linear dimension	Viscosity	Density	Thermal conductivity	Specific heat capacity at constant pressure	Temperature difference	The product of the coefficient of thermal expansion	and the acceleration due to gravity
6	7	1	ı	ď	ĸ	S	ΔT	(gg)	

It may be noted that both temperature and heat are taken as fundamental units as heat is not expressed here in terms of M, L, T.

With nine parameters and five dimensions, equation 9.55 may be rearranged in four dimensionless groups.

Using the II-theorem for solution of the equation, and taking as the recurring set: $l, \rho, \mu, \Delta T, k$

 $q, u, (\beta g), C_p$

The non-recurring variables are:

Then:

$$l \equiv L$$

$$l \equiv L$$

$$\rho \equiv ML^{-3}$$

$$\mu \equiv ML^{-1}T^{-1}$$

$$T = ML^{-1}\mu^{-1} = \rho l^{3}l^{-1}\mu^{-1} = \rho l^{2}\mu^{-1}$$

$$\Delta T \equiv \theta$$

$$\theta \equiv \Delta T$$

$$k \equiv HL^{-1}T^{-1}\theta^{-1}$$

$$H = kLT\theta = kl\rho l^{2}\mu^{-1}\Delta T = kl^{3}\rho\mu^{-1}\Delta T$$

 $\Gamma = 1$

The II groups are then:

$$\Pi_1 = q\mathbf{H}^{-1}\mathbf{L}^2\mathbf{T} = qk^{-1}l^{-3}\rho^{-1}\mu\Delta T^{-1}l^2\rho l^2\mu^{-1} = qk^{-1}l\Delta T^{-1}$$

$$\Pi_2 = uL^{-1}T = ul^{-1}\rho l^2\mu^{-1} = u\rho l\mu^{-1}$$

$$\Pi_3 = C_p \mathbf{H}^{-1} \mathbf{M} \theta = C_p k^{-1} l^{-3} \rho^{-1} \mu \Delta T^{-1} \rho l^3 \Delta T = C_p k^{-1} \mu$$

$$\Pi_4 = \beta g L^{-1} \mathbf{T}^2 \theta = \beta g l^{-1} \rho^2 l^4 \mu^{-2} \Delta T = \beta g \Delta T \rho^2 \mu^{-2} l^3$$

The relation in equation 9.55 becomes:

$$\frac{ql}{k\Delta T} = \frac{hl}{k} = \phi \left[\left(\frac{l\mu\rho}{\mu} \right) \left(\frac{C_p\mu}{k} \right) \left(\frac{\beta g\Delta Tl^3\rho^2}{\mu^2} \right) \right]$$
(9.56)

 $Nu = \phi[Re, Pr, Gr]$

This general equation involves the use of four dimensionless groups, although it may frequently be simplified for design purposes. In equation 9.56:

 u_l/k - is known as the Nusselt group Nu (already referred to in equation 9.46),

 $lu\rho/\mu$ the Reynolds group Re,

 $C_p \mu/k$ the Prandtl group Pr, and

 $\beta g \Delta T l^3 \rho^2 / \mu^2$ the *Grashof* group *Gr*

It is convenient to define other dimensionless groups which are also used in the analysis of heat transfer. These are:

$$lu\rho C_p/k$$
 the Peclet group, $Pe = RePr$,

$$GC_p/kl$$
 the Graetz group Gz , and

$$h/C_p \rho u$$
 the Stanton group, $St = Nu/(RePr)$

It may be noted that many of these dimensionless groups are ratios. For example, the Nusselt group h/(k/l) is the ratio of the actual heat transfer to that by conduction over a thickness l, whilst the Prandtl group, $(\mu/\rho)/(k/C_\rho\rho)$ is the ratio of the kinematic viscosity to the thermal diffusivity.

For conditions in which only natural convection occurs, the velocity is dependent on the buoyancy effects alone, represented by the Grashof number, and the Reynolds group may be omitted. Again, when forced convection occurs the effects of natural convection are usually negligible and the Grashof number may be omitted. Thus:

for natural convection:
$$Nu = f(Gr, Pr)$$
 (9.57)

and for forced convection:
$$Nu = f(Re, Pr)$$
 (9.58)

For most gases over a wide range of temperature and pressure, $C_p \mu/k$ is constant and the Prandtl group may often be omitted, simplifying the design equations for the calculation of film coefficients with gases.

9.4.3. Forced convection in tubes

Turbulent flow

The results of a number of workers who have used a variety of gases such as air, carbon dioxide, and steam and of others who have used liquids such as water, acetone, kerosene, and benzene have been correlated by DITTUS and BOELTER⁽¹³⁾ who used mixed units for their variables. On converting their relations using consistent (SI, for example) units, they

or heating of fluids:

$$Nu = 0.0241Re^{0.8}P_{\tau^{0.4}} \tag{9.59}$$

and for cooling of fluids:

$$Nu = 0.0264Re^{0.8}Pr^{0.3} (9.60)$$

In these equations all of the physical properties are taken at the mean bulk temperature of the fluid $(T_i + T_o)/2$, where T_i and T_o are the inlet and outlet temperatures. The difference in the value of the index for heating and cooling occurs because in the former case the film temperature will be greater than the bulk temperature and in the latter case less. Conditions in the film, particularly the viscosity of the fluid, exert an important effect on the heat transfer process.

Subsequently MCADAMS⁽¹⁴⁾ has re-examined the available experimental data and has concluded that an exponent of 0.4 for the Prandtl number is the most appropriate one for both heating and cooling. He also has slightly modified the-coefficient to 0.023 (corresponding to Colburn's value, given below in equation 9.64) and gives the following equation, which applies for Re > 2100 and for fluids of viscosities not exceeding 2 mN s/m²:

$$Nu = 0.023Re^{0.8}Pr^{0.4} (9.61)$$

WINTERTON⁽¹⁵⁾ has looked into the origins of the "Dittus and Boelter" equation and has found that there is considerable confusion in the literature concerning the origin of equation 9.61 which is generally referred to as the Dittus-Boelter equation in the literature on heat transfer.

An alternative equation which is in many ways more convenient has been proposed by Colenga and includes the Stanton number $(St = h/C_p \rho u)$ instead of the Nusselt number. This equation takes the form:

$$j_H = StPr^{0.67} = 0.023Re^{-0.2} (9.62)$$

where j_H is known as the j-factor for heat transfer.

It may be noted that:

$$\frac{h}{C_p \rho u} = \left(\frac{hd}{k}\right) \left(\frac{\mu}{ud\rho}\right) \left(\frac{k}{C_p \mu}\right)$$

$$St = NuRe^{-1}Pr^{-1} (9.63)$$

us multiplying equation 9.62 by RePr^{0.33}:

(9.64)

chais a form of equations 9.59 and 9.60.

HEAT TRANSFER

α

Writing a heat balance for the flow through a tube of diameter d and length l with a rise in temperature for the fluid from T_i to T_o :

$$h\pi dl \Delta T = \frac{\pi d^2}{4} C_p u \rho (T_o - T_i)$$

$$St = \frac{h}{C_p \rho u} = \frac{d(T_o - T_t)}{4!\Delta T}$$
 (9.65)

where ΔT is the mean temperature difference between the bulk fluid and the walls.

With very viscous liquids there is a marked difference at any position between the viscosity of the fluid adjacent to the surface and the value at the axis or at the bulk temperature of the fluid. SIEDER and TATE⁽¹⁷⁾ examined the experimental data available

and suggested that a term $\left(\frac{\mu}{\mu_J}\right)^{0.14}$ be included to account for the viscosity variation and the fact that this will have opposite effects in heating and cooling. (μ is the viscosity at the bulk temperature and μ_s the viscosity at the wall or surface). They give a logarithmic plot, but do not propose a correlating equation. However, MCADAMS⁽¹⁴⁾ gives the following equation, based on Sieder and Tate's work:

$$Nu = 0.027Re^{0.8}P_r^{0.33} \left(\frac{\mu}{\mu_s}\right)^{0.14}$$
 (9.66)

This equation may also be written in the form of the Colburn equation (9.62).

When these equations are applied to heating or cooling of gases for which the Prandtl group usually has a value of about 0.74, substitution of Pr = 0.74 in equation 9.64 gives:

$$Nu = 0.020Re^{0.8} (9.67)$$

Water is very frequently used as the cooling medium and the effect of the variation of physical properties with temperature may be included in equation 9.64 to give a simplified equation which is useful for design purposes (Section 9.9.4).

There is a very big difference in the values of h for water and air for the same linear velocity. This is shown in Figures 9.21–9.23 and Table 9.2, all of which are based on the work of FISHENDEN and SAUNDERS⁽¹⁸⁾.

The effect of length to diameter ratio (1/d) on the value of the heat transfer coefficient may be seen in Figure 9.24. It is important at low Reynolds numbers but ceases to be significant at a Reynolds number of about 10^4 .

It is also important to note that the film coefficient varies with the distance from the entrance to the tube. This is especially important at low (I/d) ratios and an average value is given approximately by: h_{average} $|A\rangle_{0.7}$

$$\frac{h_{\text{average}}}{h_{\infty}} = 1 + \left(\frac{d}{l}\right)^{0.7} \tag{9.68}$$

where h_{∞} is the limiting value for a very long tube.

The roughness of the surface of the inside of the pipe can have an important bearing on rates of heat transfer to the fluid, although COPE⁽¹⁹⁾, using degrees of artificial roughness

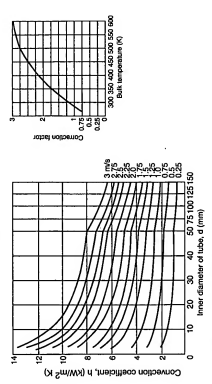


Figure 9.21. Film coefficients of convection for flow of water through a tube at 289 K

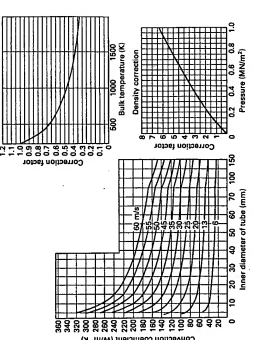
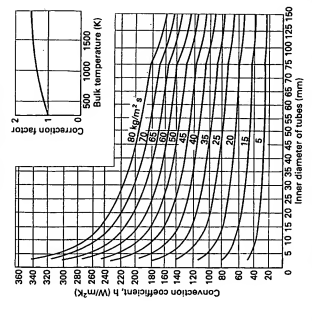


Figure 9.22. Film coefficients of convection for flow of air through a tube at various velocities (289 K, 101.3 kN/m²)





Film coefficients of convection for flow of air through a tube for various mass velocities (289 K, $101.3~{\rm kN/m^2})$ Figure 9.23.

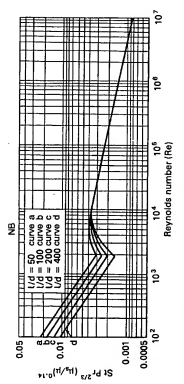


Figure 9.24. Effect of length: diameter ratio on heat transfer coefficient

per cent higher. It was concluded that, for the same pressure drop, greater heat transfer was obtained from a smooth rather than a rough tube. The effect of a given scale deposit is usually less serious for gases than water because of the higher thermal resistance of ranging from 0.022 to 0.14 of the pipe diameter, found that, although the friction loss was some six times greater than for smooth tubes, the heat transfer was only 100-120

HEAT TRANSFER

Table 9.2. Film coefficients for air and water (289 K and 101.3 kN/m²)

of tube	of tube	Vel	Velocity	Mass v	Mass velocity	Film coefficie	Film coefficient of heat transfer h
(mm)	(in)	(ш/s)	(ft/s)	(kg/m² s)	(lb/ft² h)	(W/m ² K) [Ref.18]	(Btu/h ft²°F) [Ref.18]
ir					`		
25	1.0	'n	16.4	6.11	4530	31.2	5.5
		10	32.8	12.2	9050	50.0	8.8
		20	65.6	24.5	18,100	84.0	14.8
		4	131	48.9	36,200	146	25.7
		8	197	73.4	54,300	211	37.2
20	2.0	'n	16.4	6.11	4530	23.8	4.2
		10	32.8	12.2	9050	44.9	7.9
		20	65.6	24.5	18,100	77.8	13.7
		4	131	48.9	36,200	127	22.4
		9	197	73.4	54,300	181	31.9
75	3.0	S	16.4	6.11	4530	21.6	3.8
		10	32.8	12.2	9050	39.7	7.0
		20	65.6	24.5	18,100	71.0	12.5
		9	131	48.9	36,200	119	21.0
		9	161	73.4	54,300	169	29.8
later .							
23	1.0	0.5	26.	488	361,000	2160	380
		1.0	3.28	975	722,000	3750	099
		1.5	4.92	1460	1,080,000	5250	925
		2.0	6.55	1950	1,440,000	6520	1150
		2.5	8.18	2440	1,810,000	7780	1370
23	2.0	0.5	<u>2</u> .	488	361,000	1870	330
		1.0	3.28	975	722,000	3270	575
		1.5	4.92	1460	1,080,000	4540	800
		2.0	6.55	1950	1,440,000	2590	985
		2.5	8.18	2440	1,810,000	0029	1180
75	3.0	0.5	1.6	488	361,000	1760	.310
		1.0	3.28	975	722,000	3070	5 40
		1.5	4.92	1460	1,080,000	4200	740
		2.0	6.55	1950	1,440,000	5220	920
		4	0 10	2440	1 010 000	000	51.

the gas film, although layers of dust or of materials which sublime may seriously reduce heat transfer between gas and solid by as much as 40 per cent.

Streamline flow

Although heat transfer to a fluid in streamline flow takes place solely by conduction, it is convenient to consider it here so that the results may be compared with those for turbulent flow.

In Chapter 3 it has been seen that, for streamline flow through a tube, the velocity distribution across a diameter is parabolic, as shown in Figure 9.25. If a liquid enters a section heated on the outside, the fluid near the wall will be at a higher temperature than will therefore be greater in the heated section, and correspondingly less at the centre. The velocity distribution will therefore be altered, as shown. If the fluid enters a section where it is cooled, the same reasoning will show that the distribution in velocity will be altered that in the centre and its viscosity will be lower. The velocity of the fluid near the wall

Figure 9.25. Effect of heat transfer on the velocity distribution for a liquid

to that shown. With a gas the conditions are reversed, because of the increase of viscosity with temperature. The heat transfer problem is therefore complex.

and:

for heat flow decreases with distance from the surface. The problem can conveniently be divided into two parts. Firstly, heat transfer in the entry length in which the boundary layers are limited in thickness to the radius of the pipe and, furthermore, the effective area layers are developing, and, secondly, heat transfer under conditions of fully developed For the common problem of heat transfer between a fluid and a tube wall, the boundary flow. Boundary layer flow is discussed in Chapter 11.

For the region of fully developed flow in a pipe of length L, diameter d and radius r, the rate of flow of heat Q through a cylindrical surface in the fluid at a distance y from the wall is given by:

$$Q = -k2\pi L(r - y)\frac{d\theta}{dy}$$
 (9.69)

Thus:

and:

Close to the wall, the fluid velocity is low and a negligible amount of heat is carried along the pipe by the flowing fluid in this region and Q is independent of y.

Thus:
$$\frac{d\theta}{dy} = -\frac{Q}{k2\pi L}(r - y)^{-1} \text{ and } \left(\frac{d\theta}{dy}\right)_{y=0} = -\frac{Q}{2\pi kLr}$$

$$\frac{d^2\theta}{dy^2} = -\frac{Q}{k2\pi L}(r - y)^{-2} \text{ and } \left(\frac{d^2\theta}{dy^2}\right)_{y=0} = -\frac{Q}{2\pi kLr^2}$$
Thus:
$$\left(\frac{d^2\theta}{dy^2}\right)_{y=0} = r^{-1}\left(\frac{d\theta}{dy}\right)_{y=0}$$
(9.7)

Assuming that the temperature of the walls remains constant at the datum temperature and that the temperature at any distance y from the walls is given by a polynomial, then:

(9.70)

Thus:

Thus:
$$\theta = a_0 y + b_0 y^2 + c_0 y^3$$
 (9.71) $\frac{d\theta}{dy} = a_0 + 2b_0 y + 3c_0 y^2$ and $\left(\frac{d\theta}{dy}\right)_{y=0} = a_0$

$$\frac{d^2\theta}{dy^2} = 2b_0 + 6c_0 y \quad \text{and} \quad \left(\frac{d^2\theta}{dy^2}\right)_{y=0} = 2i$$

 $2b_0 = \frac{a_0}{r} \quad \text{(from equation 9.65)}$

Thus:

$$b_0 = \frac{a_0}{2r}$$

and:

If the temperature of the fluid at the axis of the pipe is θ_s and the temperature gradient at the axis, from symmetry, is zero, then:

giving:
$$0 = a_0 + 2r \left(\frac{a_0}{2r}\right) + 3c_0 r^2$$

$$c_0 = -\frac{2a_0}{3r^2}$$
and:
$$\theta_s = a_0 r + r^2 \left(\frac{a_0}{2r}\right) + r^3 \left(\frac{-2a_0}{3r^2}\right)$$

$$= \frac{5}{6} a_0 r$$

$$a_0 = \frac{6}{5} \frac{\theta_s}{r}$$

$$a_0 = \frac{6}{5} \frac{\theta_s}{r^2}$$
and:
$$c_0 = -\frac{4}{5} \frac{\theta_s}{r^3}$$
Thus:
$$\frac{\theta}{\rho} = \frac{6}{5} \frac{\gamma}{r} + \frac{3}{2} \left(\frac{\gamma}{2}\right)^2 - \frac{4}{2} \left(\frac{\gamma}{2}\right)^3$$

Thus the rate of heat transfer per unit area at the wall:

$$q = -k \left(\frac{d\theta}{dy}\right)_{y=0}$$

$$= -\frac{6k\theta_x}{5r}$$
 (9.3)

temperature of the fluid is defined as the ratio of the heat content to the heat capacity of In general, the temperature θ_s at the axis is not known, and the heat transfer coefficient is related to the temperature difference between the walls and the bulk fluid. The bulk the fluid flowing at any section. Thus the bulk temperature θ_B is given by: $C_p \rho \theta u_x 2\pi (r-y) \, \mathrm{d}y$

$$\theta_B = \frac{\int_0^r C_p \rho \theta u_x 2\pi (r - y) \, \mathrm{d}y}{\int_0^r C_p \rho u_x 2\pi (r - y) \, \mathrm{d}y}$$

$$= \frac{\int_0^r 6u_x(r-y) \, dy}{\int_0^r u_x(r-y) \, dy}$$
(9.74)

From Poiseuille's law (equation 3.30):

$$u_x = \frac{-\Delta P}{4\mu L} [r^2 - (r - y)^2] = \frac{-\Delta P}{4\mu L} (2ry - y^2)$$

where u_s is the velocity at the pipe axis,

$$\frac{u_x}{u_c} = \frac{2y}{r} - \left(\frac{y}{r}\right)^2$$

(9.76)

Thus:
$$\int_{0}^{r} u_{x}(r-y) \, dy = r^{2} u_{s} \int_{0}^{1} \left[2\frac{y}{r} - \left(\frac{y}{r}\right)^{2} \right] \left(1 - \frac{y}{r} \right) \, d\left(\frac{y}{r}\right)$$
$$= r^{2} u_{s} \int_{0}^{1} \left[2\left(\frac{y}{r}\right) - 3\left(\frac{y}{r}\right)^{2} + \left(\frac{y}{r}\right)^{3} \right] \, d\left(\frac{y}{r}\right)$$

$$=\frac{1}{4}r^2u_s$$

(9.77)

$$\frac{\theta}{\theta_s} = \frac{6}{5} \frac{y}{r} + \frac{3}{5} \left(\frac{y}{r}\right)^2 - \frac{4}{5} \left(\frac{y}{r}\right)^3$$
 (equation 9.72)

$$\int_{0}^{r} \theta u_{r}(r-y) \, \mathrm{d}y = r^{2} u_{s} \theta_{s} \int_{0}^{1} \left[\frac{6}{5} \frac{y}{r} + \frac{3}{5} \left(\frac{y}{r} \right)^{2} - \frac{4}{5} \left(\frac{y}{r} \right)^{3} \right] \left[2 \left(\frac{y}{r} \right) - 3 \left(\frac{y}{r} \right)^{2} + \left(\frac{y}{r} \right)^{3} \right] \mathrm{d} \left(\frac{y}{r} \right)$$

$$= r^2 u_s \theta_s \int_0^1 \left[\frac{12}{5} \left(\frac{y}{r} \right)^2 - \frac{12}{5} \left(\frac{y}{r} \right)^3 - \frac{11}{5} \left(\frac{y}{r} \right)^4 + 3 \left(\frac{y}{r} \right)^5 - \frac{4}{5} \left(\frac{y}{r} \right)^6 \right] d \left(\frac{y}{r} \right)$$

$$= r^2 u_s \theta_s \left(\frac{4}{5} - \frac{3}{5} - \frac{11}{25} + \frac{1}{2} - \frac{4}{35} \right)$$

$$= \frac{51}{5} r^2 u_s \theta_s$$

Substituting from equations 9.77 and 9.78 in equation 9.74:

9.11 and 9.18 in equation
$$\theta_B = \frac{\frac{51}{350}r^2u_s\theta_s}{\frac{1}{4}r^2u_s}$$

$$= \frac{192}{175}\theta_s = 0.583\theta_s$$

(9.79)

The heat transfer coefficient h is then given by:

$$h = -\frac{q}{\theta_r}$$

where q is the rate of heat transfer per unit area of tube.

Thus, from equations 9.73 and 9.79:

$$h = \frac{6k\theta_s/5r}{0.583\theta_s} = \frac{2.06k}{r} = 4.1\frac{k}{d}$$

$$h = \frac{0.583\theta_s}{0.583\theta_s} = \frac{-r}{r} = 4.1\frac{-r}{d}$$

$$Nu = \frac{hd}{r} = 4.1$$

This expression is applicable only to the region of fully developed flow. The heat transfer coefficient for the inlet length can be calculated approximately, using the expressions given in Chapter 11 for the development of the boundary layers for the flow over a plane surface. It should be borne in mind that it has been assumed throughout that the physical properties of the fluid are not appreciably dependent on temperature and therefore the expressions will not be expected to hold accurately if the temperature differences are large and if the properties vary widely with temperature.

For values of (RePr d/I) greater than 12, the following empirical equation is applicable:

(9.75)

$$Nu = 1.62 \left(RePr \frac{d}{l} \right)^{1/3} = 1.75 \left(\frac{GC_p}{kl} \right)^{1/3}$$
 (9.81)

where $G = (\pi d^2/4)\rho u$, i.e. the mass rate of flow. The product RePr is termed the Peclet number Pe.

$$Pe = \frac{ud\rho}{\mu} \frac{Cp\mu}{k} = \frac{Cp\rho ud}{k} \tag{9.3}$$

Equation 9.81 may then be written:

$$Nu = 1.62 \left(Pe \frac{d}{l} \right)^{1/3} \tag{9.83}$$

In this equation the temperature difference is taken as the arithmetic mean of the terminal values, that is:

$$(T_w - T_1) + (T_w - T_2)$$

where T_w is the temperature of the tube wall which is taken as constant. If the liquid is heated almost to the wall temperature T_w (that is when GC_p/kl is very small) then, on equating the heat gained by the liquid to that transferred from the pipe:

$$GC_p(T_2 - T_1) = \pi dlh \frac{T_2 - T_1}{2}$$

(9.78)

$$h = \frac{2}{\pi} \frac{GC_p}{dl} \tag{9}$$

For values of (RePrd/l) less than about 17, the Nusselt group becomes approximately constant at 4.1; the value given in equation 9.80. Experimental values of h for viscous oils are greater than those given by equation 9.81 for heating and less for cooling. This is due to the large variation of viscosity with temperature and the correction introduced for turbulent flow may also be used here, giving:

$$Nu\left(\frac{\mu_s}{\mu}\right)^{0.14} = 1.86\left(RePr\frac{d}{l}\right)^{1/3} = 2.01\left(\frac{GC_p}{kl}\right)^{1/3} \tag{9.85}$$

or:
$$Nu\left(\frac{\mu_s}{\mu}\right)^{0.14} = 1.86\left(Pe\frac{d}{l}\right)^{1/3}$$
 (9.86)

important. The equations are not entirely consistent since for very small values of ΔT the equation 9.84 applies. These equations have been obtained with tubes about 10 mm to 40 mm in diameter, and the length of unheated tube preceding the heated section is constants in equations 9.81 and 9.85 would be expected to be the same. It is important When $(GC_p/kl) < 10$, the outlet temperature closely approaches that of the wall and to note, when using these equations for design purposes, that the error may be as much as ±25 per cent for turbulent flow and greater for streamline conditions.

With laminar flow there is a marked influence of tube length and the curves shown in Figure 9.24 show the parameter l/d from 50 to 400.

Whenever possible, streamline conditions of flow are avoided in heat exchangers because of the very low heat transfer coefficients which are obtained. With very viscous Since the viscosity of these materials is critically dependent on temperature, the equations liquids, however, turbulent conditions can be produced only if a very high pressure drop across the plant is permissible. In the processing industries, streamline flow in heat exchangers is most commonly experienced with heavy oils and brines at low temperatures. would not be expected to apply with a high degree of accuracy.

9.4.4. Forced convection outside tubes

Flow across single cylinders

across which air is passed is not uniform but is a maximum at the front and rear, and a minimum at the sides, where the rate is only some 40 per cent of the maximum. The If a fluid passes at right angles across a single tube, the distribution of velocity around the tube will not be uniform. In the same way the rate of heat flow around a hot pipe general picture is shown in Figure 9.26 but for design purposes reference is made to the

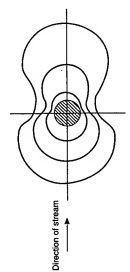


Figure 9.26. Distribution of the film heat transfer coefficient round a cylinder with flow normal to the axis for three different values of Re

have studied the flow of a hot gas past a single cylinder, varying from a thin wire to a A number of workers, including, REIHER⁽²⁰⁾, HILPERT⁽²¹⁾, GRIFFITHS and AWBERY⁽²²⁾,

HEAT TRANSFER

tube of 150 mm diameter. Temperatures up to 1073 K and air velocities up to 30 m/s have been used with Reynolds numbers $(d_o u \rho/\mu)$ from 1000 to 100,000 (where d_o is the cylinder diameter, or the outside tube diameter). The data obtained may be expressed

$$Nu = 0.26Re^{0.6}Pr^{0.3} (9.87)$$

Taking Pr as 0.74 for gases, this reduces to

$$Nu = 0.24Re^{0.6} (9.8)$$

DAVIS⁽²³⁾ has also worked with water, paraffin, and light oils and obtained similar results. For very low values of Re (from 0.2 to 200) with liquids the data are better represented by the equation:

$$Nu = 0.86Re^{0.43} Pr^{0.3} (9.89)$$

In each case the physical properties of the fluid are measured at the mean film temperature T_f , taken as the average of the surface temperature T_w and the mean fluid temperature T_m ; where $T_m = (T_1 + T_2)/2$.

Flow at right angles to tube bundles

changing. Moreover the degree of turbulence is considerably less for banks of tubes in line, as at (a), than for staggered tubes, as at (b) in Figure 9.27. With the small bundles which are common in the processing industries, the selection of the true mean area for One of the great difficulties with this geometry is that the area for flow is continually flow is further complicated by the change in number of tubes in the rows.

The results of a number of workers for heat transfer to and from gases flowing across tube banks may be expressed by the equation:

$$Nu = 0.33C_h Re_{\text{max}}^{0.6} Pr^{0.3} \tag{9.90}$$

GRIMISON⁽²⁴⁾ proposed this form of expression to correlate the data of HUGE⁽²⁵⁾ and PIERSON(26) who worked with small electrically heated tubes in rows of ten deep. Other workers have used similar equations. Some correction factors have been given by PIERSON(26) for bundles with less than ten rows although there are insufficient reported data from commercial exchangers to fix these values with accuracy. Thus for five rows a where C_h depends on the geometrical arrangement of the tubes, as shown in Table 9.3. factor of 0.92 and for eight rows 0.97 is suggested.

These equations are based on the maximum velocity through the bundle. Thus for an in-line arrangement as is shown in Figure 9.27a, $G'_{\text{max}} = G'Y/(Y - d_o)$, where Y is the pitch of the pipes at right-angles to direction of flow; it is more convenient here to use the mass flowrate per unit area G' in place of velocity. For staggered arrangements the maximum velocity may be based on the distance between the tubes in a horizontal line or on the diagonal of the tube bundle, whichever is the less.

It has been suggested that, for in-line arrangements, the constant in equation 9.90 should be reduced to 0.26, but there is insufficient evidence from commercial exchangers to confirm this.

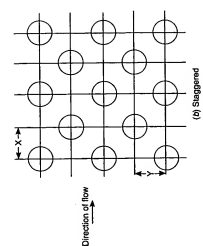


Figure 9.27. Arrangements of tubes in heat exchangers

Table 9.3.(18) Values of Ch and Cf

		. ~	1	28	1.74	20		80	1.16	96
	Staggered	c_f		2.5	=	1.		7	-	Ö
.5do	Sta	C,		1.16	1.05	1.02		1.15	1.02	0.98
$X=1.5d_o$	In-line	C_f		1.74	1.56	1.32		0.97	96.0	. 0.85
	l-aī	Ch		1.06	1.00	1.00		0.95	96.0	96.0
	paras	C_f	$Y = 1.25d_o$	2.52	1.56	1.26	$Y = 1.5d_o$	1.80	1.10	0.88
.25do	Staggered	์	*	1.21	1.06	1.03		1.17	1.08	0.99
$X = 1.25d_o$	ine	C,		1.68	4	1.20		0.79	0.84	0.74
	In-line	ర		1.06	1.00	1.00		0.95	96.0	96.0
		Remax		2000	20.000	40,000		2000	20,000	40,000

With liquids the same equation may be used, although for Re less than 2000, there is insufficient published work to justify an equation. MCADAMS, $^{(27)}$ however, has given a curve for h for a bundle with staggered tubes ten rows deep. An alternative approach has been suggested by KERN $^{(28)}$ who worked in terms of the

hydraulic mean diameter de for flow parallel to the tubes:

HEAT TRANSFER

429

 $d_e = 4 \times \frac{\text{Free area for flow}}{\text{Wetted perimeter}}$

.:. :::

$$=4\left(\frac{Y^2-(\pi d_o^2/4)}{-3}\right)$$

for a square pitch as shown in Figure 9.28. The maximum cross-flow area A_s is then given by:

$$A_s = \frac{d_s l_B C'}{\gamma}$$

where C' is the clearance, l_B the baffle spacing, and d_s the internal diameter of the shell.

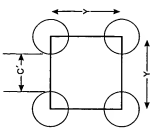


Figure 9.28. Clearance and pitch for tube layouts

The mass rate of flow per unit area G'_2 is then given as rate of flow divided by A_3 , and the film coefficient is obtained from a Nusselt type expression of the form:

$$\frac{h_0 d_e}{k} = 0.36 \left(\frac{d_e G_s'}{\mu} \right)^{0.55} \left(\frac{C_p \mu}{k} \right)^{1/3} \left(\frac{\mu}{\mu_s} \right)^{0.14}$$
 (9.91)

There are insufficient published data to assess the relative merits of equations 9.90 and 9.91

For 19 mm tubes on 25 mm square pitch:

$$d_e = 4 \frac{[25^2 - (\pi/4)19^2]}{\pi \times 19}$$
$$= 22.8 \text{ mm or } 0.023 \text{ m}$$

Example 9.9

14.4 tonnefh (4.0 kg/s) of nitrobenzene is to be cooled from 400 to 315 K by heating a stream of benzene from 305 to 345 K.

Two tubular heat exchangers are available each with a 0.44 m i.d. shell fitted with 166 tubes, 19.0 mm o.d. and 15.0 mm i.d., each 5.0 m long. The tubes are arranged in two passes on 25 mm square pitch with a baffle spacing of 150 mm. There are two passes on the shell side and operation is to be countercurrent. With benzene passing through the tubes, the anticipated film coefficient on the tube side is 1000 W/m²K.

Assuming true cross-flow prevails in the shell, what value of scale resistance could be allowed if these units

For nitrobenzene: $C_p = 2380$ J/kg K, k = 0.15 W/m K, $\mu = 0.70$ mN s/m²

(i) Tube side coefficient.

$$h_i = 1000 \text{ W/m}^2 \text{ K}$$
 based on inside area
$$\frac{1000 \times 15.0}{19.0} = 790 \text{ W/m}^2 \text{ K}$$
 based on outside area
$$\frac{19.0}{19.0}$$

(ii) Shell side coefficient.

.: 0

Area for flow = shell diameter \times baffle spacing \times clearance/pitch

$$= \frac{0.44 \times 0.150 \times 0.006}{0.025} = 0.0158 \text{ m}^2$$

$$=\frac{0.44 \times 0.130 \times 0.000}{0.025} = 0.$$

 $G_{\rm r}' = \frac{4.0}{0.0158} = 253.2 \text{ kg/m}^2 \text{s}$

Taking $\mu/\mu_s = 1$ in equation 9.91:

$$h_o = 0.36 \frac{k}{d_e} \left(\frac{d_e G_s'}{\mu} \right)^{0.55} \left(\frac{C_p \mu}{k} \right)^{0.33}$$

The hydraulic mean diameter,

$$d_e = 4\left[\left(25^2 - \frac{\pi \times 19.0^2}{4}\right) / (\pi \times 19.0)\right] = 22.8 \text{ mm} \text{ or } 0.023 \text{ m}$$

$$h_o = \left(\frac{0.15}{0.023}\right) 0.36 \left(\frac{0.023 \times 253.2}{0.70 \times 10^{-3}}\right)^{0.55} \left(\frac{2380 \times 0.70 \times 10^{-3}}{0.15}\right)^{0.33}$$

 $= 2.35 \times 143 \times 2.23 = 750 \text{ W/m}^2 \text{ K}$

The logarithmic mean temperature difference is given by:

$$\Delta T_m = \frac{(400 - 345) - (315 - 305)}{\ln(400 - 345)/(315 - 305)}$$

= 26.4 deg K

The corrected mean temperature difference is then $\Delta T_m \times F = 26.4 \times 0.8 = 21.1$ deg K (Details of the correction factor for ΔT_m are given in Section 9.9.3)

 $Q = 4.0 \times 2380(400 - 315) = 8.09 \times 10^5 \text{ W}$

The surface area of each tube $= 0.0598 \text{ m}^2/\text{m}$

$$U_o = \frac{Q}{A_o \Delta T_m F} = \frac{8.09 \times 10^5}{2 \times 166 \times 5.0 \times 0.0598 \times 21.1}$$
= 386.2 W/m² K

If scale resistance is R_d , then:

$$R_d = \frac{1}{386.2} - \frac{1}{750} - \frac{1}{1000} = \underline{0.00026 \text{ m}^2 \text{ K/W}}$$

This is a rather low value, though the heat exchangers would probably be used for this duty.

HEAT TRANSFER

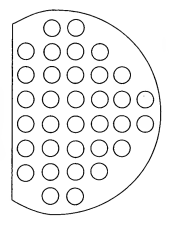


Figure 9.29. Baffle for heat exchanger

Reference may also be made in Volume 6 and to the work of SHORT⁽²⁹⁾, DONOHUE⁽³⁰⁾, and TINKER⁽³¹⁾. The various methods are all concerned with developing a method of As discussed in Section 9.9 it is common practice to fit baffles across the tube bundle in order to increase the velocity over the tubes. The commonest form of baffle is shown in Figure 9.29 where it is seen that the cut-away section is about 25 per cent of the total area. With such an arrangement, the flow pattern becomes more complex and the extent of leakage between the tubes and the baffle, and between the baffle and the inside of the shell of the exchanger, complicates the problem, as discussed further in Section 9.9.6. calculating the true area of flow and of assessing the probable influence of leaks. When using baffles, the value of h_o , as found from equation 9.89, is commonly multiplied by 0.6 to allow for leakage although more accurate approaches have been developed as discussed in Section 9.9.6.

The drop in pressure for the flow of a fluid across a tube bundle may be important because of the small pressure head available and because by good design it is possible to get a better heat transfer for the same drop in pressure. $-\Delta P_f$ depends on the velocity u_i through the minimum area of flow and in Chapter 3 an equation proposed by GRIMISON⁽²⁴⁾

$$-\Delta P_f = \frac{C_{fj}\rho u_t^2}{6}$$
 (equation 3.83)

Table 9.4.⁽¹⁸⁾ Ratio of heat transfer to friction for tube bundles ($Re_{max} = 20,000$)

		$X=1.25d_o$	10		$X = 1.5d_o$. 0
	ζ	5	C_h/C_f	ರೆ	6,7	C_h/C_f
In-line						
$Y = 1.25d_o$	1	<u>4</u> .	0.69	_	1.56	9.0
$Y = 1.5d_o$	96.0	0.84	1.14	96.0	0.96	1.0
Staggered						
$Y = 1.25d_o$	1.06	1.56	0.68	1.05	1.74	09.0
$Y = 1.5d_o$	<u>2</u> .	1.10	0.95	1.02	1.16	0.88

where C_f depends on the geometry of the tube layout and j is the number of rows of tubes. It is found that the ratio of C_h , the heat transfer factor in equation 9.90, to C_f

is greater for the in-line arrangement but that the actual heat transfer is greater for the staggered arrangement, as shown in Table 9.4.

The drop in pressure $-\Delta P_f$ over the tube bundles of a heat exchanger is also given by:

$$-\Delta P_f = \frac{f'G_s^2(n+1)d_v}{2pd_s}$$
 (9.92)

ë.

where f' is the friction factor given in Figure 9.30, G'_s the mass velocity through bundle, n the number of baffles in the unit, d_v the inside shell diameter, ρ the density of fluid, d_e the equivalent diameter, and $-\Delta P_f$ the drop in pressure.

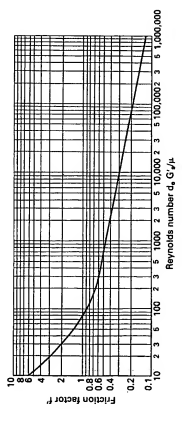


Figure 9.30. Friction factor for flow over tube bundles

Example 9:10

54 tonneth (15 kg/s) of benzene is cooled by passing the stream through the shell side of a tubular heat exchanger, 1 m i.d., fitted with 5 m tubes, 19 mm o.d. arranged on a 25 mm square pitch with 6 mm olearance. If the baffle spacing is 0.25 m (19 baffles), what will be the pressure drop over the tube bundle? $(\mu = 0.5 \text{ mN s/m}^2)$.

Solution

Cross-flow area:
$$A_{\rm J} = \frac{1.0 \times 0.25 \times 0.006}{0.005} = 0.06 \, \text{m}^2$$

$$G_{s}' = \frac{15}{0.06} = 250 \text{ kg/m}^2 \text{ s}$$

 $d_{s} = \frac{4[0.025^2 - (\pi/4)0.019^2]}{(\pi/4)0.019^2} = 0.0229 \text{ m}$

 $\pi \times 0.019$

Equivalent diameter:

Mass flow:

Reynolds number through the tube bundle =
$$\frac{250 \times 0.0229}{0.5 \times 10^{-3}} = 11450$$

From Figure 9.29:
$$f' = 0.280$$

$$= 881 \text{ kg/m}^3$$

Density of benzene From Figure 9.29:

HEAT TRANSFER

From equation 9.92:

$$-\Delta P_f = \frac{0.280 \times 250^2 \times 20 \times 1.0}{2 \times 881 \times 0.0229} = \frac{8674 \text{ N/m}^2}{8674}$$

$$\frac{8674}{881 \times 9.81} = \frac{1.00 \text{ m of benzene}}{1.00 \text{ m of benzene}}$$

9.4.5. Flow in non-circular sections

Rectangular ducts

For the heat transfer for fluids flowing in non-circular ducts, such as rectangular ventilating ducts, the equations developed for turbulent flow inside a circular pipe may be used if an equivalent diameter, such as the hydraulic mean diameter de discussed previously, is used in place of d.

The data for heating and cooling water in turbulent flow in rectangular ducts are reasonably well correlated by the use of equation 9.59 in the form:

$$\frac{hd_e}{k} = 0.023 \left(\frac{d_e G'}{\mu} \right)^{0.8} \left(\frac{C_p \mu}{k} \right)^{0.4} \tag{9.93}$$

Whilst the experimental data of COPE and BAILEY(32) are somewhat low, the data of WASHINGTON and MARKS⁽³³⁾ for heating air in ducts are well represented by this equation.

Annular sections between concentric tubes

Concentric tube heat exchangers are widely used because of their simplicity of construction and the ease with which additions may be made to increase the area. They also give turbulent conditions at low volumetric flowrates.

in selecting the best equivalent diameter to use. When considering the film on the outside In presenting equations for the film coefficient in the annulus, one of the difficulties is of the inner tube, DAVIS(34) has proposed the equation:

$$\frac{hd_1}{k} = 0.031 \left(\frac{d_1 G'}{\mu} \right)^{0.8} \left(\frac{C_p \mu}{k} \right)^{0.33} \left(\frac{\mu}{\mu_s} \right)^{0.14} \left(\frac{d_2}{d_1} \right)^{0.15}$$
 (9.94)

where d_1 and d_2 are the outer diameter of the inner tube, and the inner diameter of the outer tube, respectively.

CARPENTER et $al.^{(35)}$ suggest using the hydraulic mean diameter $d_e = (d_2 - d_1)$ in the Sieder and Tate equation (9.66) and recommend the equation:

$$\frac{hd_e}{k} \left(\frac{\mu_s}{\mu}\right)^{0.14} = 0.027 \left(\frac{d_e G'}{\mu}\right)^{0.8} \left(\frac{C_p \mu}{k}\right)^{0.33} \tag{9.95}$$

Their data, which were obtained using a small annulus, are somewhat below those given by equation 9.95 for values of d_eG'/μ less than 10,000, although this may be because the flow was not fully turbulent: with an index on the Reynolds group of 0.9, the equation fitted the data much better. There is little to choose between these two equations, but they both give rather high values for h.

For the viscous region, Carpenter's results are reasonably well correlated by the equa-

$$\frac{hd_e}{k} \left(\frac{\mu_s}{\mu}\right)^{0.14} = 2.01 \left(\frac{GC_p}{kl}\right)^{0.33}$$

$$= 1.86 \left[\left(\frac{d_e G'}{\mu}\right) \left(\frac{C_p \mu}{k}\right) \left(\frac{d_1 + d_2}{l}\right) \right]^{1/3}$$
(9.97)

Equations 9.96 and 9.97 are the same as equations 9.85 and 9.86, with d_{ϵ} replacing d. These results have all been obtained with small units and mainly with water as the fluid in the annulus.

Flow over flat plates

For the turbulent flow of a fluid over a flat plate the Colburn type of equation may be used with a different constant:

$$j_h = 0.037 Re_x^{-0.2} \tag{9.98}$$

where the physical properties are taken as for equation 9.64 and the characteristic dimension in the Reynolds group is the actual distance x along the plate. This equation therefore gives a point value for j_h .

9.4.6. Convection to spherical particles

In Section 9.3.4, consideration is given to the problem of heat transfer by conduction through a surrounding fluid to spherical particles or droplets. Relative motion between the fluid and particle or droplet causes an increase in heat transfer, much of which may be due to convection. Many investigators have correlated their data in the form:

$$Nu' = 2 + \beta'' Re'^n Pr^m \tag{9.99}$$

where values of eta'', a numerical constant, and exponents n and m are found by experiment. In this equation, Nu' = hd/k and $Re' = du\rho/\mu$, the Reynolds number for the particle, uis the relative velocity between particle and fluid, and d is the particle diameter. As the relative velocity approaches zero, Re' tends to zero and the equation reduces to Nu' = 2for pure conduction.

provided further experimental data, have concluded that for particle Reynolds numbers in RowE et al. (36), having analysed a large number of previous studies in this area and the range 20-2000, equation 9.99 may be written as:

$$Nu' = 2.0 + \beta'' Re^{0.5} Pr^{0.33}$$
 (9 100)

where β'' lies between 0.4 and 0.8 and has a value of 0.69 for air and 0.79 for water. In some practical situations the relative velocity between particle and fluid may change due

tion (equation 10.233) applies, with the Sherwood number replacing the Nusselt number to particle acceleration or deceleration, and the value of Nu' can then be time-dependent. For mass transfer, which is considered in more detail in Chapter 10, an analogous relaand the Schmidt number replacing the Prandtl number.

9.4.7. Natural convection

HEAT TRANSFER

If a beaker containing water rests on a hot plate, the water at the bottom of the beaker becomes hotter than that at the top. Since the density of the hot water is lower than that of the cold, the water in the bottom rises and heat is transferred by natural convection. In the air near the surface being hotter and of lower density than that some distance away. In the transfer of heat occurs at a correspondingly lower rate since the natural convection the same way air in contact with a hot plate will be heated by natural convection currents, both of these cases there is no external agency providing forced convection currents, and currents move rather slowly.

For these processes which depend on buoyancy effects, the rate of heat transfer might be expected to follow a relation of the form:

$$Nu = f(Gr, Pr)$$
 (equation 9.57)

Measurements by SCHMIDT⁽³⁷⁾ of the upward air velocity near a 300 mm vertical plate show that the velocity rises rapidly to a maximum at a distance of about 2 mm from the olate and then falls rapidly. However, the temperature evens out at about 10 mm from the plate. Temperature measurements around horizontal cylinders have been made by RAY⁽³⁸⁾.

Natural convection from horizontal surfaces to air, nitrogen, hydrogen, and carbon dioxide, and to liquids (including water, aniline, carbon tetrachloride, glycerol) has been studied by several workers, including DAVIS⁽³⁹⁾, ACKERMANN⁽⁴⁰⁾, FISHENDEN and SAUNDERS⁽¹⁸⁾ and SAUNDERS⁽⁴¹⁾. Most of the results are for thin wires and tubes up to about 50 mm diameter; the temperature differences used are up to about 1100 deg K with gases and about 85 deg K with liquids. The general form of the results is shown in Figure 9.31, where $\log Nu$ is plotted against $\log (Pr Gr)$ for streamline conditions. The curve can be represented by a relation of the form:

$$Nu = C'(Gr Pr)^n (9.101)$$

Numerical values of C' and n, determined experimentally for various geometries, are given in Table 9.5⁽⁴²⁾. Values of coefficients may then be predicted using the equation:

$$\frac{hl}{k} = C' \left(\frac{\beta g \Delta T l^3 \rho^2}{\mu^2} \frac{C_p \mu}{k} \right)^n \quad \text{or} \quad h = C' \left(\frac{\Delta T}{l} \right)^n k \left(\frac{\beta g \rho^2 C_p}{\mu k} \right)^n \tag{9.10}$$

Table 9.5. Values of C', C'' and n for use in equations 9.102 and 9.105⁽⁴²⁾

Geometry	GrPr	ن	u	C" (SI units) (for air at 294 K)
Vertical surfaces	< 104	1.36	0.20	
(l = vertical dimension < 1 m)	104-109	0.59	0.25	1.37
	> 109	0.13	0.33	1.24
Horizontal cylinders				
(l = diameter < 0.2 m)	$10^{-5} - 10^{-3}$	0.71	90.0	
	$10^{-3} - 1.0$	1.09	0.10	
	$1.0 - 10^4$	1.09	0.20	
	$10^4 - 10^9$	0.53	0.25	1.32
	> 109	0.13	0.33	1.24
Horizontal flat surfaces				
(facing upwards)	$10^5 - 2 \times 10^7$	0.54	0.25	1.86
(facing upwards)	$2 \times 10^7 - 3 \times 10^{10}$	0.14	0.33	
(facing downwards)	$3 \times 10^5 - 3 \times 10^{10}$	0.27	0.25	0.88

2.2 2.0 8 9. 4. 5 0. 9.9 9.0 0.4 0.2

HEAT TRANSFER

Example 9.11

Estimate the heat transfer coefficient for natural convection from a horizontal pipe 0.15 m diameter, with a surface temperature of 400 K to air at 294 K

Over a wide range of temperature, $k^4(\beta g \rho^2 C_P / \mu k) = 36.0$ For air at a mean temperature of 0.5(400 + 294) = 347 K, k = 0.0310 W/m K (Table 6, Appendix A1)

$$\frac{\beta g \rho^2 C_p}{\mu k} = \frac{36.0}{0.0310^4} = 3.9 \times 10^7$$

Thus:

% 80 80

$$GrPr = 3.9 \times 10^7 (400 - 294) \times 0.15^3$$

$$= 1.39 \times 10^7$$

$$n = 0.25$$
 and $C'' = 1.32$

$$h = 1.32(400 - 294)^{0.25} (0.15)^{(3 \times 0.25) - 1}$$

For the transfer of heat from a hot surface across a thin layer of fluid to a parallel cold

$$\frac{Q}{Q_k} = \frac{h\Delta T}{(k/x)\Delta T} = \frac{hx}{k} = Nu \tag{9.106}$$

for gases, the coefficient of cubical expansion β is taken as 1/T, where T is the absolute

where the physical properties are at the mean of the surface and bulk temperatures and,

Figure 9.31. Natural convection from horizontal tubes

2 3 log (Gr Pr)

٦

Ŋ ግ 4

φ

For vertical plates and cylinders, KATO et al. (43) have proposed the following equations

for situations where 1 < Pr < 40:

 $Nu = 0.138Gr^{0.36}(Pr^{0.175} - 0.55)$

For $Gr > 10^9$:

where Q_k is the rate at which heat would be transferred by pure thermal conduction

between the layers, a distance x apart, and Q is the actual rate. For $(Gr\ Pr) = 10^3$, the heat transferred is approximately equal to that due to conduction alone, though for $10^4 < Gr Pr < 10^6$, the heat transferred is given by:

(9.103)

(9.104)

 $\left(\frac{Pr}{0.861 + Pr}\right)$

 $Nu = 0.683Gr^{0.25}Pr^{0.25}$

and for $Gr < 10^9$:

Simplified dimensional equations have been derived for air, water and organic liquids by grouping the fluid properties into a single factor in a rearrangement of equation 9.102

Natural convection to air

$$\frac{Q}{Q_k} = 0.15(Gr\ Pr)^{0.25} \tag{9.107}$$

independent of surface area, provided that the linear dimensions of the surfaces are large compared with x. For higher values of $(Gr\ Pr)$, Q/Q_k is proportional to $(Gr\ Pr)^{1/3}$, which is noted in Figure 9.32. In this equation the characteristic dimension to be used for the Grashof group is x, the distance between the planes, and the heat transfer is showing that the heat transferred is not entirely by convection and is not influenced by the distance x between the surfaces.

concentric cylinders. It is important to note from this general analysis that a single layer A similar form of analysis has been given by KRAUSSOLD⁽⁴⁴⁾ for air between two of air will not be a good insulator because convection currents set in before it becomes 25 mm thick. The good insulating properties of porous materials are attributable to the

(9.105)

Values of C" (in SI units) are also given in Table 9.5 for air at 294 K. Typical values for

water and organic liquids are 127 and 59 respectively.

 $h = C''(\Delta T)^n l^{3n-1} (W/m^2 K)$

Solution

From Equation 9.102:

800

uM gol

From Table 9.5:

Thus, in Equation 9.104:

$$h = 1.32(400 - 294)^{0.25}(0.15)^{0.25}$$
$$= 1.32 \times 106^{0.25} \times 0.15^{-0.25}$$
$$= \frac{6.81 \text{ W/m}^2 \text{ K}}{6.81 \text{ W/m}^2 \text{ K}}$$

Fluids between two surfaces

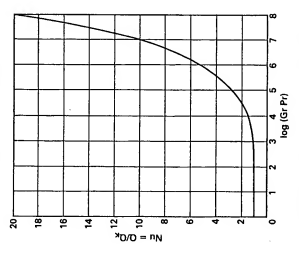


Figure 9.32. Natural convection between surfaces

fact that they offer a series of very thin layers of air in which convection currents are not present.

9.5. HEAT TRANSFER BY RADIATION

9.5.1. Introduction

It has been seen that heat transfer by conduction takes place through either a solid or a natural movement of a hot fluid. The third mechanism of heat transfer, radiation, can take many fluids are transparent to radiation, and it is generally assumed that the emission of thermal radiation is by "waves" of wavelengths in the range $0.1-100 \mu$ m which travel in straight lines. This means that direct radiation transfer, which is the result of an interchange stationary fluid and heat transfer by convection takes place as a result of either forced or place without either a solid or a fluid being present, that is through a vacuum, although between various radiating bodies or surfaces, will take place only if a straight line can be drawn between the two surfaces; a situation which is often expressed in terms of one surface "seeing" another. Having said this, it should be noted that opaque surfaces sometimes cast shadows which inhibit radiation exchange and that indirect transfer by radiation can take place as a result of partial reflection from other surfaces. Although all bodies at temperatures in excess of absolute zero radiate energy in all directions, radiation is of especial importance from bodies at high temperatures such as those encountered in furnaces, boilers and high temperature reactors, where in addition to radiation from hot surfaces, radiation from reacting flame gases may also be a consideration.

HEAT TRANSFER

9.5.2. Radiation from a black body

In thermal radiation, a so-called *black body* absorbs all the radiation falling upon it, regardless of wavelength and direction, and, for a given temperature and wavelength, no surface can emit more energy than a black body. The radiation emitted by a black body, whilst a function of wavelength and temperature, is regarded as *diffuse*, that is, it is independent of direction. In general, most rough surfaces and indeed most engineering materials may be regarded as being diffuse. A black body, because it is a perfect emitter or absorber, provides a standard against which the radiation properties of real surfaces may be compared.

If the emissive power E of a radiation source—that is the energy emitted per unit area per unit time—is expressed in terms of the radiation of a single wavelength λ , then this is known as the monochromatic or spectral emissive power E_{λ} , defined as that rate at which radiation of a particular wavelength λ is emitted per unit surface area, per unit wavelength in all directions. For a black body at temperature T, the spectral emissive power of a wavelength λ is given by Planck's Distribution Law:

$$E_{\lambda,b} = C_1/[\lambda^3(\exp(C_2/\lambda T) - 1)]$$
 (9.108)

where, in SI units, $E_{\lambda,b}$ is in W/m³ and $C_1 = 3.742 \times 10^{-16}$ W/m² and $C_2 = 1.439 \times 10^{-2}$ mK are the respective radiation constants. Equation 9.108 permits the evaluation of the emissive power from a black body for a given wavelength and absolute temperature and values obtained from the equation are plotted in Figure 9.33 which is based on the work of INCROPERA and DE WITT⁽⁴⁵⁾. It may be noted that, at a given wavelength, the radiation from a black body increases with temperature and that, in general, short wavelengths are associated with high temperature sources.

Example 9.12

What is the temperature of a surface coated with carbon black if the emissive power at a wavelength of 1.0×10^{-6} m is 1.0×10^9 W/m³? How would this be affected by a +2 per cent error in the emissive power measurement?

Solution

From equation 9.108 $\exp(C_2/\lambda T) = [C_1/E_{\lambda,b}\lambda^5) + 1]$

or: $\exp(1.439 \times 10^{-2}/(1.0 \times 10^{-6}T)) = [3.742 \times 10^{-16}/(1 \times 10^{-9} \times (1.0 \times 10^{-6})^{5})]$

 $= 3.742 \times 10^5$

 $(1.439 \times 10^4)/T = \ln(3.742 \times 10^5) = 12.83$

Thus:

and

 $T = (1.439 \times 10^4)/12.83 = 1121 \text{ K}$

With an error of +2 per cent, the correct value is given by:

 $E_{\lambda,b} = (100 - 2)(1 \times 10^9)/100 = 9.8 \times 10^8 \text{ W/m}^3$

 $9.8 \times 10^{-8} = (3.742 \times 10^{-16})/[(1 \times 10^{-6})^{5}]$

fin equation 9.108:

 $(\exp(1.439\times 10^{-2}/(1.0\times 10^{-6}T))-1)]$

T = 1120 K

and:

Thus, the error in the calculated temperature of the surface is only 1 K.

temperature by Wein's Displacement Law, which states that the wavelength for maximum The wavelength at which maximum emission takes place is related to the absolute emission varies inversely with the absolute temperature of the source, or:

$$\lambda_{max} T = \text{constant}, C_3 (= 2.898 \times 10^{-3} \text{ mK in SI units})$$
 (9.109)

Thus, combining equations 9.108 and 9.109:

$$E_{\lambda max,b} = C_1 / [(C_3/T)^5 [\exp(C_2/C_3) - 1]]$$

 $E_{\lambda max,b} = C_4 T^5$ (9.110)

ö

where, in SI units, the fourth radiation constant, $C_4 = 12.86 \times 10^{-6} \text{ W/m}^3 \text{ K}^5$. Values of the maximum emissive power are shown by the broken line in Figure 9.33.

only a limited range of wavelengths. This warms the air in the greenhouse, though to a the wavelength will be much longer, as seen from Figure 9.33, and one at which the glass will not transmit radiation. In this way, radiation outwards from within a greenhouse is much lower temperature than that of the external source, the sun; a temperature at which considerably reduced and the air within the enclosure retains its heat. Much the same An interesting feature of Figure 9.33 is that it illustrates the well-known greenhouse effect which depends on the ability of glass to transmit radiation from a hot source over

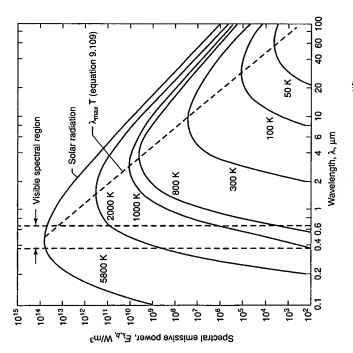


Figure 9.33. Spectral black-body emissive power (45)

HEAT TRANSFER

phenomenon occurs in the gases above the earth's surface which will transmit incoming ation through gases and indeed through glass is one example of a situation where the radiation from the sun at a given wavelength though not radiation from the earth which, because it is at a lower temperature, emits at a longer wavelength. The passage of raditransmissivity t, discussed in Section 9.5.3, is not zero.

unit time per unit area of surface over all wavelengths and in all directions. This may be mining the area corresponding to a particular temperature under the Planck distribution The total emissive power E is defined as the rate at which radiation energy is emitted per determined by making a summation of all the radiation at all wavelengths, that is by detercurve, Figure 9.33. In this way, from equation 9.108, the total emissive power is given by:

$$E_b = \int_0^\infty C_1 \, d\lambda / [\lambda^5 (\exp(C_2/\lambda T) - 1)]$$
 (9.111)

which is known as the Stefan-Boltzmann Law. This may be integrated for any constant value of T to give:

$$E_b = \sigma T^4 \tag{9.112}$$

where, in SI units, the Stefan-Boltzmann constant $\sigma = 5.67 \times 10^{-8} \; \mathrm{W/m^2 \; K^4}$

Example 9.13

Electrically-heated carbide elements, 10 mm in diameter and 0.5 m long, radiating essentially as black bodies, are to be used in the construction of a heater in which thermal radiation from the surroundings is negligible. If the surface temperature of the carbide is limited to 1750 K, how many elements are required to provide a radiated thermal output of 500 kW?

Solution

From equation 9.112, the total emissive power is given by:

$$E_b = \sigma T^4 = (5.67 \times 10^{-8} \times 1750^4) = 5.32 \times 10^5 \text{ W/m}^2$$

The area of one element = $\pi(10/1000)0.5 = 1.571 \times 10^{-2} \text{ m}^2$

Power dissipated by one element = $(5.32\times10^5\times1.571\times10^{-2})$ = 8.367×10^3 W and:

Number of elements required = $(500 \times 1000)/(8.357 \times 10^3) = 59.8$ say $\frac{60}{2}$ Thus:

9.5.3. Radiation from real surfaces

wavelength) to that emitted by a black body at the same temperature. Emissivities of The emissivity of a material is defined as the ratio of the radiation per unit area emitted from a "real" or from a grey surface (one for which the emissitivity is independent of roughness of the material, and possibly on the wavelength and direction of the emitted radiation as well. For diffuse surfaces where emissivities are independent of direction, the "real" materials are always less than unity and they depend on the type, condition and emissivity, which represents an average over all directions, is known as the hemispherical emissivity. For a particular wavelength λ this is given by:

$$\mathbf{e}_{\lambda} = E_{\lambda} E_b \tag{9.113}$$

and, similarly, the total hemispherical emissivity, an average over all wavelengths, is given by:

$$\mathbf{e} = E/E_b \tag{9.114}$$

of incident radiation absorbed, and the emissivity of a surface are equal. If two bodies A and B of areas A_1 and A_2 are in a large enclosure from which no energy is lost, then the energy absorbed by A from the enclosure is A₁a₁I where I is the rate at which energy is falling on unit area of A and a_1 is the absorptivity. The energy given out by A is E_1A_1 Equation 9.114 leads to Kirchoff's Law which states that the absorptivity, or fraction and, at equilibrium, these two quantities will be equal or:

$$IA_1\mathbf{a}_1 = A_1E_1$$

 $IA_2\mathbf{a}_2 = A_2E_2$ and, for B:

 $E_1/a_1 = E_2/a_2 = E/a$ for any other body.

Since $E/a = E_b/a_b$, then, from equation 9.114:

$$e = E/E_b = a/a_b$$

and, as by definition, $a_b = 1$, the emissivity of any body is equal to its absorptivity, or:

(9.115)

For most industrial, non-metallic surfaces and for non-polished metals, e is usually about 0.9. although values as low as 0.03 are more usual for highly polished metals such

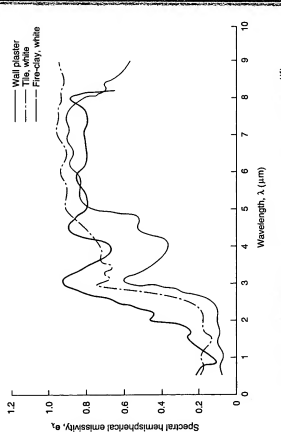


Figure 9.34. Spectral emissivity of non-conductors as a function of wavelength (45)

HEAT TRANSFER

as copper or aluminium. As explained later, a small cavity in a body acts essentially as a black body with an effective emissivity of unity. The variation of emissivity with is based on the work of HOTTEL and SAROFIM(46). More complete data are available in wavelength is illustrated in Figure 9.34⁽⁴⁵⁾ and typical values are given in Table 9.6 which Appendix A1, Table 10. If equation 9.113 is written as:

$$E_{\lambda} = \mathbf{e}_{\lambda} E_{\lambda,b} \tag{9.1}$$

As shown in Figure 9.35, for a temperature of 2000 K for example, the emission curve for a real material may have a complex shape because of the variation of emissivity with wavelength, If, however, the ordinate of the black body curve $E_{\lambda,b}$ at a particular wavelength is multiplied by the spectral emissivity of the source at that wavelength, the then the spectral emissive power of a grey surface may be obtained from the spectral emissivity, e_{λ} and the spectral emissive power of a black body $E_{\lambda,b}$ given by equation 9.108. ordinates on the curve for the real surface are obtained, and the total emissive power of he real surface is obtained by integrating E_{λ} over all possible wavelengths to give:

$$E = \int_0^\infty E_\lambda \, d\lambda = \int_0^\infty \mathbf{e}_\lambda E_{\lambda,b} \, d\lambda \tag{9.117}$$

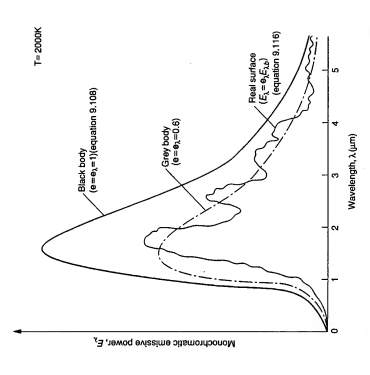


Figure 9.35. Comparison of black body, grey body and real surface radiation at 2000 K. (45)

This integration may be carried out numerically or graphically, though this approach, which has been considered in some detail by INCROPERA and DE WITT⁽⁴⁵⁾, can be difficult, especially where the spectral distribution of radiation arrives at a surface of complex structure. The amount of calculation involved cannot often be justified in practical situations and it is more usual to use a mean spectral emissivity for the surface which is assumed to be constant over a range of wavelengths. Where the spectral emissivity does not vary with wavelength then the surface is known as a grey body, and, for a diffuse grey body, e, can be replaced by e in equations 9.112 and 9.114

$$E = \mathbf{e}E_b = \mathbf{e}\sigma T^4 \tag{9.118}$$

In this way, the emissive power of a grey body is a constant proportion of the power emitted by the black body, resulting in the curve shown in Figure 9.35 where, for example, e = 0.6. The assumption that the surface behaves as a grey body is valid for most engineering calculations if the value of emissivity is taken as that for the dominant temperature of the radiation.

From equation 9.117, it is seen that the rate of heat transfer by radiation from a hot body at temperature T_1 to a cooler one at temperature T_2 is then given by:

$$q = Q/A = e\sigma(T_1^4 - T_2^4) = e\sigma(T_1 - T_2)(T_1^3 + T_1^2T_2 + T_1T_2^2 + T_2^3)$$

The quantity $q/(T_1 - T_2)$ is a heat transfer coefficient as used in convective heat transfer, and here it may be designated h_r , the heat transfer coefficient for radiation heat transfer where:

$$h_r = q/(T_1 - T_2) = \frac{e\sigma(T_1^4 - T_2^4)}{T_1 - T_2} = e\sigma(T_1^3 + T_1^2T_2 + T_1T_2^2 + T_3^3)$$
(9.119)

It may be noted that if $(T_1 - T_2)$ is very small, that is T_1 and T_2 are virtually equal, then:

$$h_r = 4e\sigma T^3$$

Example 9.14

What is the emissivity of a grey surface, 10 m² in area, which radiates 1000 kW at 1500 K? What would be the effect of increasing the temperature to 1600 K?

Solution

The emissive power $E = (1000 \times 1000)/10 = 100,000 \text{ W/m}^2$ From equation 9.118: $e = E/\sigma T^4$ $= 100,000/(5.67 \times 10^{-8} \times 1500^4) = 0.348$

 $E = e\sigma T^4$ = (0.348 × 5.67 × 10⁻⁸ × 1600⁴) = 1295 kW

At 1600 K:

an increase of 29.5 per cent for a 100 deg K increase in temperature.

HEAT TRANSFER

Table 9.6. Typical emissivity values (46)

soling and metallic oxides	rides		
(A) Metats and metatic o.	_	ì	•
Aluminium	Polished plate	296	0.040
	Kougn plate	567	0.055
Brass	Folished	311-589	0.096
Copper	Polished	390	0.023
77.0	Flate, oxidised	498	0.78
Cold	ruginy pousied	200-200	0.018-0.35
Iron and steel	Polished fron	700-1300	0.144-0.377
	Cast iron, newly turned	295	0.435
	Smooth sheet tron	1172-1311	0.55-0.60
	Sheet steel, oxidised	295	0.657
	Iron	373	0.736
	Steel plate, rough	311-644	0.94-0.97
Lead	Pure, unoxidised	400-500	0.057-0.075
	Grey, oxidised	297	0.281
Mercury		273-373	0.09-0.12
Molybdenum	Filament	1000-2866	0.096-0.292
Monel	Metal oxidised	472-872	0.41-0.46
Nickel	Polished	200-600	0.07-0.087
	Wire	460-1280	0.096-0.186
	Plate, oxidised	472-872	0.37-0.48
Nickel alloys	Chromonickel	325-1308	0.64-0.76
	Nickelin, grey oxidised	294	0.262
Platinum	Pure, polished plate	200-000	0.054-0.104
	Strip	1200-1900	0.12-0.17
	Filament	300-1600	0.036-0.192
	Wire ·	200-1600	0.073-0.182
Silver	Polished	310-644	0.0221-0.0312
Fantalum	Filament	1600-3272	0.194 - 0.31
Tin	Bright tinned iron sheet	298	0.043-0.064
Tungsten	Filament	3588	0.39
Zinc	Pure, polished	200-009	0.045-0.053
	Galvanised sheet	297	0.276
(B) Refractories, building materials, paints etc.	naterials, paints etc.		
Asbestos	Board	297	96.0
Brick	Red, rough	294	0.93
	Silica, unglazed	1275	0.80
Carbon	Filament	1311-1677	0.526
	Candle soot	372-544	0.952
	Lampblack	311-644	0.945
Enamel	White fused on iron	292	0.897
Glass	Smooth	295	0.937
Paints, lacquers,	Snow-white enamel	596	9060
	Black, shiny lacquer	298	0.875
	Black matt shellac	350-420	0.91
Plaster,	Lime, rough	283-361	0.91
Porcellain,	Glazed	295	0.924
Refractory materials	Poor radiators	872-1272	0.65-0.75
	Good radiators	872-1272	0.80-0.90
Rubber	Hard, glossy plate	296	0.945
	Soft, grey, rough	298	0.859
Water			

In a real situation, radiation incident upon a surface may be absorbed, reflected and transmitted and the properties, absorptivity, reflectivity and transmissivity may be used to describe this behaviour. In theory, these three properties will vary with the direction and wavelength of the incident radiation, although, with diffuse surfaces, directional variations may be ignored and mean, hemispherical properties used.

If the absorptivity a, the fraction of the incident radiation absorbed by the body is defined on a spectral basis, then:

$$\mathbf{a}_{\lambda} = I_{\lambda, abs}/I_{\lambda} \tag{9.120}$$

and the total absorptivity, the mean over all wavelengths, is defined as:

$$\mathbf{a} = I_{abs}/I \tag{9.121}$$

Since a black body absorbs all incident radiation then for a black body:

$$a_{\lambda} = a = 1$$

The absorptivity of a grey body is therefore less than unity.

In a similar way, the reflectivity, r, the fraction of incident radiation which is reflected from the surface, is defined as:

$$\mathbf{r} = I_{\text{ref}}/I \tag{9.122}$$

and the transmissivity, t, the fraction of incident radiation which is transmitted through

$$t = I_{\rm trans}/I \tag{9.123}$$

Since, as shown in Figure 9.36, all the incident radiation is absorbed, reflected or transmitted, then:

$$I_{abs} + I_{ref} + I_{trans} = I$$
$$a + r + t = 1$$

ä

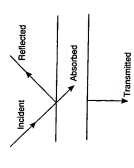


Figure 9.36. Reflection, absorption, and transmission of radiation

Since most solids are opaque to thermal radiation, t = 0 and therefore:

$$+\mathbf{r} = 1 \tag{9.124}$$

HEAT TRANSFER

Kirchoff's Law, discussed previously, states that, at any wavelength, the emissivity and the absorptivity are equal. If this is extended to total properties, then, at a given temperature:

$$= a$$
 (equation 9.115)

a small aperture approximates to a black body in that radiation entering is subjected to For a grey body, the emissivity and the absorptivity are, by definition, independent where one radiation property (a, r or e) is specified for an opaque body, the other two may be obtained from equations 9.115 and 9.124. Kirchoff's Law explains why a cavity with repeated internal absorption and reflection so that only a negligible amount of the incident radiation escapes through the aperture. In this way, $\mathbf{a} = \mathbf{e} = 1$ and, at T K, the emissive of temperature and hence equation 9.115 may be applied more generally showing that, power of the aperture is σT^4 .

9.5.4. Radiation transfer between black surfaces

therefore only a proportion of the radiation leaving a surface arrives at any other given surface. This proportion depends on the relative geometry of the surfaces and this may be Since radiation arriving at a black surface is completely absorbed, no problems arise from multiple reflections. Radiation is emitted from a diffuse surface in all directions and taken into account by the view factor, shape factor or configuration F, which is normally written as F_{ij} for radiation arriving at surface j from surface i. In this way, F_{ij} , which is, of course, completely independent of the surface temperature, is the fraction of radiation leaving i which is directly intercepted by j.

If radiant heat transfer is taking place between two black surfaces, 1 and 2, then:

radiation emitted by surface
$$1 = A_1 E_{b_1}$$

The fraction of this radiation which arrives at and is totally absorbed by surface 2 is F_{12} where A_1 and E_{b1} are the area and black body emissive power of surface 1, respectively. and the heat transferred is then:

$$Q_{1\rightarrow 2} = A_1 F_{12} E_{b1}$$

Similarly, the radiation leaving surface 2 which arrives at 1 is given by:

$$Q_{2\to 1} = A_2 F_{21} E_{b2}$$

and the net radiation transfer between the two surfaces is $Q_{12}=(Q_{1\rightarrow 2}-Q_{2\rightarrow 1})$ or:

$$Q_{12} = A_1 F_{12} E_{b1} - A_2 F_{21} E_{b2}$$

= $\sigma A_1 F_{12} T_1^4 - \sigma A_2 F_{21} T_2^4$ (9.1)

When the two surfaces are at the same temperature, $T_1 = T_2$, $Q_{12} = 0$

$$Q_{12} = 0 = \sigma T_1^4 (A_1 F_{12} - A_2 F_{21})$$

and thus:

Since the temperature T_1 can have any value so that, in general $T_1 \neq 0$, then:

$$A_1 F_{12} = A_2 F_{21} \tag{9.12}$$

HEAT TRANSFER

$$Q_{12} = \sigma A_1 F_{12}(T_1^4 - T_2^4) = \sigma A_2 F_{12}(T_1^4 - T_2^4)$$
 (9.127)

The product of an area and an appropriate view factor is known as the exchange area which, in SI units, is expressed in m2. In this way, A1F12 is known as exchange area 1-2.

Example 9.15

Calculate the view factor, F_{21} and the net radiation transfer between two black surfaces, a rectangle 2 m by 1 m (area A_1) at 1500 K and a disc 1 m in diameter (area A_2) at 750 K, if the view factor, $F_{12} = 0.25$.

Solution

$$A_1 = (2 \times 1) = 2 \text{ m}^2$$

$$A_2 = (\pi \times 1^2)/4 = 0.785 \text{ m}^2$$

$$A_2 = (\pi \times 1^2)/4 = 0.78$$

From equation 9.126:
$$A_1F_{12} = A_2F_{21}$$

$$(2 \times 0.25) = 0.785F_{21}$$

$$F_{21} = 0.637$$

and:

In equation 9.127:
$$Q_{12} = \sigma A_1 F_{12}(T_1^4 - T_2^4)$$

$$Q_{12} = (5.67 \times 10^{-8} \times 2 \times 0.25)(1500^4 - 750^4)$$

$$= 5.38 \times 10^5$$
 W or 538 kW

View factors, the values of which determine heat transfer rates, are dependent on the geometrical configuration of each particular system. As a simple example, radiation may be considered between elemental areas dA_1 and dA_2 of two irregular-shaped flat bodies, well separated by a distance L between their mid-points as shown in Figure 9.37. If α_1 and α_2 are the angles between the imaginary line joining the mid-points and the normals, the rate of heat transfer is then given by:

$$Q_{12} = \sigma(T_1^4 - T_2^4) \int^{A_1} \int^{A_2} (\cos \alpha_1 \cos \alpha_2 \, dA_1 \, dA_2) / \pi L^2$$
 (9.128)

Equation 9.128 may be extended to much larger surfaces by subdividing these into a series of smaller elements, each of exchange area $A_i F_{ij}$, and summing the exchange areas between each pair of elements to give:

$$A_i F_{ij} = A_j F_{ji} = \int_{A_i} \int_{A_j} (\cos \alpha_i \cos \alpha_j \, dA_i \, dA_j)/\pi L^2$$
 (9.129)

In this procedure, the value of the integrand can be determined numerically for every pair of elements and the double integral, approximately the sum of these values, then becomes:

$$\int_{A_j} \int_{A_i} (\cos \alpha_i \cos \alpha_j \, dA_i \, dA_j) / \pi L^2 = \sum_{A_i} \sum_{A_j} (\cos \alpha_i \cos \alpha_j \, dA_i \, dA_j) / \pi L^2 \qquad (9.130)$$

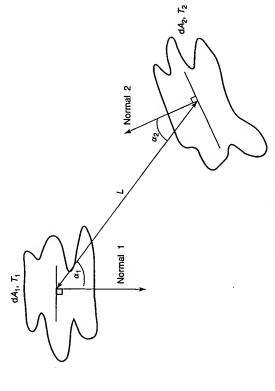


Figure 9.37. Determination of view factor

The amount of calculation involved here can be very considerable and use of a computer is usually required. A simpler approach is to make use of the many expressions, graphs and tables available in the heat transfer literature. Typical data, presented by INCROPERA and DE WITT⁽⁴⁵⁾ and by HOWELL⁽⁴⁷⁾, are shown in Figures 9.38-9.40, where it will be seen all the radiation leaving one surface arrives at the second surface as, for example, when that in many cases, the values of the view factors approach unity. This means that nearly a sphere is contained within a second larger sphere. Wherever a view factor approaches zero, only a negligible part of one surface can be seen by the other surface.

It is important to note here that if an element does not radiate directly to any part of its own surface, the shape factor with respect to itself, F11, F12 and so on, is zero. This applies to any convex surface for which, therefore, $F_{11} = 0$.

Example 9.16

What are the view factors, F_{12} and F_{21} , for (a) a vertical plate, 3 m high by 4 m long, positioned at right angles to one edge of a second, horizontal plate, 4 m wide and 6 m long, and (b) a 1 m diameter sphere positioned within a 2 m diameter sphere?

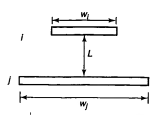
Solution

(a) Using the nomenclature in Figure 9.40 iii:

$$Y/X = (6/4) = 1.5$$
 and $Z/X = (3/4) = 0.75$

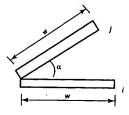
 $F_{12} = 0.12$

(i) Parallel plates with mid-lines connected by perpendicular

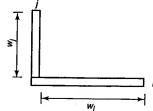


 $F_{ij} = \{[(W_i + W_j)^2 + 4]^{0.5} - [(W_j - W_i)^2 + 4]^{0.5}\}/2W_i$ $F_{ij} = 1 - \sin(\alpha/2)$ where: $W_i = W_i/L$ and $W_i = W_i/L$

(ii) Inclined parallel plates of equal width and a common edge

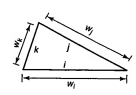


(iii) Perpendicular plates with a common edge



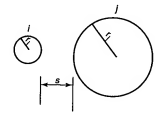
 $F_{ij} = \{1 + (w_j/w_i) - [1 + (w_j/w_i)^2]^{0.5}\}/2$

(iv) Three-sided enclosure



 $F_{ij} = (w_i + w_j - w_k)/2w_i$

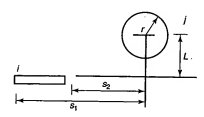
(v) Parallel cylinders of different radius



 $F_{ij} = (1/2\pi)\{-\pi + [C^2 - (R+1)^2]^{0.5} - [C^2 - (R-1)^2]^{0.5}$

+ (R-1)cos⁻¹[(R/C)-(1/C)]-(R+1)cos⁻¹[(R/C)+(1/C)]}

(vi) Cylinder and parallel rectangle



 $F_{ij} = [r/(s_1 - s_2)][\tan^{-1}(s_1/L) - \tan^{-1}(s_2/L)]$

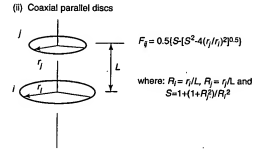
where: $R = r_i/r_i$, $S = s/r_i$ and C = 1+R+S

Figure 9.38. View Factors for two-dimensional geometries. (45)

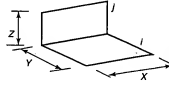
(i) Aligned parallel rectangles



 $\mathsf{F}_{\#} = [2/(\pi \bar{X} \bar{Y})] \{ \mathsf{In}[(1 + \bar{X}^2)(1 + \bar{Y}^2)/(1 + \bar{X}^2 + \bar{Y}^2)]^{0.5}$ + $\bar{X}(1+\bar{Y}^2)^{0.5} \tan^{-1} [\bar{X}/(1+\bar{Y}^2)^{0.5}]$ + $\bar{Y}(1+\bar{X}^2)^{0.5} \tan^{-1} [(\bar{Y}/(1+\bar{X}^2)^{0.5})-\bar{X} \tan^{-1} \bar{X}-\bar{Y} \tan^{-1} \bar{Y}]$ where: $\bar{X} = X/L$ and $\bar{Y} = Y/L$



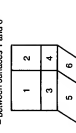
(iii) Perpendicular rectangles with a common edge



 $F_{ij} = (1/\pi W)\{W \tan^{-1}(1/W) + H \tan^{-1}(1/H) - (H^2 + W^2)^{0.5} \tan^{-1}(H^2 + W^2)^{-0.5}$ $+\ 0.25 in[(1+W^2)(1+H^2)/(1+W^2+H^2)][W^2(1+W^2+H^2)/(1+W^2)(W^2+H^2)]W^2$ $\times [H^2(1+H^2+W^2)/(1+H^2)(H^2+W^2)]H^2]$

where: H=Z/X and W=Y/X

Figure 9.39. View factors for three-dimensional geometries (45)



+ A6F6(2+4) - A5F5(1+3) - (1/2A6)(A(3+4)F(3+4×5+6) $F_{16} = (A_6/A_1)[(1/2A_6)(A_{(1+2+3+4)}F_{(1+2+3+4)(5+6)}$ - A6F6A - A5F53)]

(ii) Two parallel rectangles

8.0 8.0 p.0 X\Z

(iii) Perpendicular rectangles with a common edge

- between surfaces 1 and 7

Þ.0

+ A3F37 + A4F48] - (1/4A1)[A(1+2)F(1+2)(5+6)+A(1+4)F(1+4)(5+8) $+ A_{(2+4)} F_{(2+4)(7+8)} + A_{(2+3)} F_{(2+3)(6+7)}]$

 $F_{17} = (1/4A_1)[A_{(1+2+3+4)}F_{(1+2+3+4)(5+6+7+8)} + A_1F_{15} + A_2F_{26}$

$$\begin{split} F_{23} &= (A_{(1+2)}/A_2)[F_{(1+2)(3+4)} \cdot F_{(1+2)4}] \\ &+ (A_1/A_2)[F_{1(3+4)} \cdot F_{14}] \end{split}$$

(iii) Two parallel circular rings – between surfaces 2 and 3

the inner wall of the tube of radius x_3 and surface 1, the upper surface of the disc of radius x_1 . (iv) A circular tube and a disc between surface 3,

S.0

8.0

1.0 2 345

 $F_{13} = F_{12} \cdot F_{14}$ $F_{31} = (x_3^2/x_1^2)(F_{12} + F_{14})$

Figure 9.41. View factors obtained by using the summation rule⁽⁴⁷⁾

70.0 20.0 40.0 60.0 20.0

1.0 P

7.0 2.0 4.0 6.0 2.0

Figure 9.40. View factors for three-dimensional geometries (45)

(ii) Co-extel paratlel discs

CHEMICAL ENGINEERING

$$A_1F_{12} = A_2F_{21}$$

(3 × 4)0.12 = (4 × 6) F_{21}

and:

 $F_{21} = 0.06$

(b) For the two spheres:

$$F_{12} = 1$$
 and $F_{21} = (r_1/r_2)^2 = (1/2)^2 = \underbrace{0.25}_{======}$
 $F_{22} = 1 - (r_1/r_2)^2 = 1 - 0.25 = \underbrace{0.75}_{======}$

For a given geometry, view factors are related to each other, one example being the reciprocity relationship given in equation 9.126. Another important relationship is the summation rule which may be applied to the surfaces of a complete enclosure. In this case, all the radiation leaving one surface, say i, must arrive at all other surfaces in the enclosure so that, for n surfaces:

$$F_{11} + F_{12} + F_{13} + \cdots + F_{1n} = 1$$

 $F_{ij} = 1$

 $A_i F_{ij} = A_i$

from which:

ö

This means that the sum of the exchange areas associated with a surface in an enclosure must be same as the area of that surface. The principle of the summation rule may be extended to other geometries such as, for example, radiation from a vertical rectangle (area 1) to an adjacent horizontal rectangle (area 2), as shown in Figure 9.40iii, where they are joined to a second horizontal rectangle of the same width (area 3). In effect area 3 is an extension of area 2 but has a different view factor.

$$A_1F_{1(2+3)} = A_1F_{12} + A_1F_{13}$$

$$A_1 F_{13} = A_1 F_{1(2+3)} - A_1 F_{12} \tag{9.1}$$

ö:

Equation 9.133 allows F_{13} to be determined from the view factors F_{12} and $F_{1(2+3)}$ which can be obtained directly from Figure 9.40iii. Typical data obtained by using this technique are shown in Figure 9.41 which is based on the work of HOWELL (47).

What is the view factor F_{23} for the two parallel rings shown in Figure 9.41iii if the inner and outer radii of the two rings are: upper = 0.2 m and 0.3 m; lower = 0.3 m and 0.4 m and the rings are 0.2 m apart?

Solution

From Figure 9.41iii:

$$F_{23} = (A_{(1+2)}/A_2)(F_{(1+2)(3+4)} - F_{(1+2)4}) - (A_1/A_2)(F_{1(3+4)} - F_{14})$$

HEAT TRANSFER

Laying out the data in tabular form and obtaining F from Figure 9.40ii, then;

$$A_{(1+2)}/A_2 = 0.4^2/(0.4^2 - 0.3^2) = 2.29$$

$$A_1/A_2 = 0.3^2/(0.4^2 - 0.3^2) = 1.29$$

$$F_{23} = 2.29(0.40 - 0.22) + 1.29(0.55 - 0.30) = 0.74$$

transfer from a surface in an enclosure. If the enclosure contains n black surfaces, then Equation 9.127 may be extended in order to determine the net rate of radiation heat the net heat transfer by radiation to surface i is given by:

$$Q_i = Q_{1i} + Q_{2i} + Q_{3i} + \cdots + Q_{ni}$$

$$Q_{i} = \sum_{i=1}^{j=n} \sigma A_{j} F_{ji} (T_{j}^{4} - T_{i}^{4})$$

ö

(9.131)(9.132)

or, applying the reciprocity relationship:

$$Q_i = \sum_{j=1}^{j=n} \sigma A_i F_{ij} (T_j^4 - T_i^4)$$
 (9.134)

Example 9.18

diameter at 1200 K; the distance between the plate and the bottom of the dome being 0.5 m, as shown in Figure 9.42. If the surroundings are maintained at 290 K, the surfaces may be regarded as black bodies and heat transfer from the underside of the plate is negligible, what is the net rate of heat transfer by radiation to A plate, 1 m in diameter at 750 K, is to be heated by placing it beneath a hemispherical dome of the same

Solution

Taking area 1 as that of the plate, area 2 as the underside of the hemisphere, area 3 as an imaginary cylindrical surface linking the plate and the underside of the dome which represents the black surroundings and area 4 as an imaginary disc sealing the hemisphere and parallel to the plate, then, from equation 9.134, the net radiation to the surface of the plate 1 is given by:

$$Q_1 = \sigma A_2 F_{21}(T_2^4 - T_1^4) + \sigma A_3 F_{31}(T_3^4 - T_1^4)$$

or, using the reciprocity rule:

$$Q_1 = \sigma A_1 F_{12} (T_2^4 - T_1^4) + \sigma A_1 F_{13} (T_3^4 - T_1^4)$$

All radiation from the disc 1 to the dome 2 is intercepted by the imaginary disc 4 and hence $F_{12} = F_{14}$, which may be obtained from Figure 9.39ii, with i and j representing areas 1 and 4 respectively. Thus:

$$R_1 = r_1/L = (0.5/0.5) = 1$$
; $R_4 = r_4/L = (0.5/0.5) = 1$

and:
$$S = 1 + (1 + R_4^2)/(R_1^2) = 1 + (1 + 1.0)/(1.0) = 3.0$$

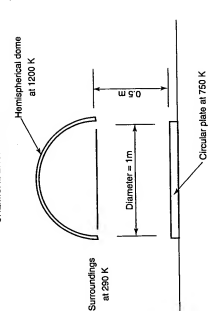


Figure 9.42. Data for Example 9.18

Thus:
$$F_{14} = 0.5 \left[S - \left[(S^2 - 4(r_4/r_1)^2]^{0.5} \right] = 0.5 \left[3 - \left[(3^2 - 4(0.5/0.5)^2]^{0.5} \right] = 0.38 \right]$$

 $F_{12} = F_{14} = 0.38$ and: The summation rule states that:

 $F_{11} + F_{12} + F_{13} = 1$

and since, for a plane surface, $F_{11} = 0$, then:

$$F_{13} = (1 - 0.38) = 0.62$$

 $A_1 = (\pi 1.0^2)/4 = 0.785 \text{ m}^2$

and hence:

$$Q_1 = (5.67 \times 10^{-8} \times 0.785 \times 0.38)(1200^4 - 750^4) + (5.67 \times 10^{-8} \times 0.785 \times 0.62)(290^4 - 750^4)$$

$$= (1.691 \times 10^{-8} \times 1.757 \times 10^{12}) - (2.760 \times 10^{-8} \times 3.093 \times 10^{11})$$

$$= 2.12 \times 10^4 \text{ W} = \frac{21.2 \text{ kW}}{2.12 \times 10^4}$$

 A_1 at temperature T_1 is radiating to a second surface 2 of area A_2 at temperature T_2 joined to it as shown in Figure 9.39iii, then adding a further surface R consisting of insulating material so as to form a triangular enclosure will reduce the heat transfer to the surroundings considerably. Even though some heat will be conducted through the Radiation between two black surfaces may be increased considerably by introducing a third surface which acts in effect as a re-radiator. For example, if a surface 1 of area insulation, this will usually be small and most of the energy absorbed by the insulated surface will be re-radiated back into the enclosure.

The net rate of heat transfer to surface 2 is given by:

$$Q_2 = \sigma A_1 F_{12} (T_1^4 - T_2^4) + \sigma A_R F_{R2} (T_R^4 - T_2^4)$$
 (9.135)

where T_R is the mean temperature of the insulation, though, in practice, there will be a temperature distribution across this surface. At steady-state, the net rate of radiation to

HEAT TRANSFER

surface R is equal to the heat loss from it to the surroundings, Q_{sur} , or:

$$Q_{\text{surr}} = \sigma A_1 F_{1R} (T_1^4 - T_R^4) + \sigma A_2 F_{2R} (T_2^4 - T_R^4)$$
 (9.136)

If Q_{sur} is negligible, that is, the surface may be treated as adiabatic, then from equation 9.136:

$$\sigma A_1 F_{1R}(T_1^4 - T_R^4) + \sigma A_2 F_{2R}(T_2^4 - T_R^4) = 0$$

Rearranging:

$$T_R^4 = (A_1 F_{1R} T_1^4 + A_2 F_{2R} T_2^4)/(A_1 F_{1R} + A_2 F_{2R})$$

Substituting for T_R from this equation in equation 9.135 and noting, from the reciprocity relationship, that $A_2F_{2R} = A_RF_{R2}$, then:

$$Q_2 = \sigma(T_1^4 - T_2^4) \{A_1 F_{12} + \{(1/(A_1 F_{1R}) + (1/(A_R F_{R2}))]^{-1}\}$$
 (9.137)

Example 9.19

A flat-bottomed cylindrical vessel, 2 m in diameter, containing boiling water at 373 K, is mounted on a cylindrical section of insulating material, 1 m deep and 2 m ID at the base of which is a radiant heater, also 2 m in diameter, with a surface temperature of 1500 K. If the vessel base and the heater surfaces may be regarded as black bodies and conduction though the insulation is negligible, what is the rate of radiant heat transfer to the vessel? How would this be affected if the insulation were removed so that the system was open to the surroundings at 290 K?

Solution

radiant heater surface and area 2 the under-surface of the vessel, with R the insulated If area 1 is the r cylinder, then:

$$A_1 = A_2 = (\pi \times 2^2/4) = 3.14 \text{ m}^2$$

$$A_R = (\pi \times 2.0 \times 1.0) = 6.28 \text{ m}^2$$

and:

From Figure 9.40ii, with i = 1, j = 2, $r_i = 1.0$ m, $r_j = 1.0$ m and L = 1.0 m,

$$(L/r_i) = (1.0/1.0) = 1.0;$$
 and $(r_j/L) = (1.0/1.0) = 1.0$

 $F_{12} = 0.40$

and:

[The view factor may also be obtained from Figure 9.39ii as follows: Using the nomenclature of Figure 9.39:

$$R_1 = (r_1/L) = (1.0/1.0) = 1.0$$

$$R_2 = (r_2/L) = (1.0/1.0) = 1.0$$

$$S = 1 + [(1 + R_2^2)/R_1^2] = 1 + [(1 + 1^2)/1^2] = 3.0$$

$$F_{12} = 0.5[S - [S^2 - 4(r_2/r_1)^2]^{0.5}] = 0.5[3 - [3^2 - (4 \times 1^2)]^{0.5}] = 0.382]$$

The summation rule states that:

and:

$$F_{11} + F_{12} + F_{1R} = 1$$

and since, for a plane surface, $F_{11} = 0$, then: $F_{1R} = (1 - 0.382) = 0.618$

Since $A_1 = A_2$:

$$F_{21} = F_{12}$$
 and $F_{2R} = F_{1R} = 0.618$

Also $A_R F_{R2} = A_2 F_{2R}$ and hence, from equation 9.137:

$$Q_2 = [A_1 F_{12} + ((1/A_1 F_{1R}) + (1/A_2 F_{2R}))^{-1} J_0(T_1^4 - T_2^4)$$

$$= \left[(3.14 \times 0.382) + ((1/3.14 \times 0.618) + (1/(3.14 \times 0.618))]^{-1} \right] (5.67 \times 10^{-8}) ((1500^4 - 373^4))$$

$$= 6.205 \times 10^5 \text{ W or } \frac{620 \text{ kW}}{620 \text{ kW}}$$

the surroundings without insulation are surface 3 at $T_3 = 290$ K, then $F_{23} = F_{2R} = 0.618$ and, from equation 9.135:

$$Q_2 = \sigma A_1 F_{12} (T_1^4 - T_2^4) + \sigma A_2 F_{23} (T_3^4 - T_2^4)$$

$$= (5.67 \times 10^{-8} \times 3.14 \times 0.382) (1500^4 - 373^4) + (5.67 \times 10^{-8} \times 3.14 \times 0.618) (290^4 - 373^4)$$

$$= 3.42 \times 10^5 \text{ W or } \frac{342 \text{ kW}}{342 \text{ kW}}; \text{ a reduction of } 45 \text{ per cent.}$$

9.5.5 Radiation transfer between grey surfaces

Since the absorptivity of a grey surface is less than unity, not all the incident radiation is with grey surfaces, reflectivity as well as the geometrical configuration must be taken into account. With grey bodies, it is convenient to consider the total radiation leaving a surface Q_o , that is the emitted plus the reflected components. The equivalent flux, $Q_o/A = Q_o$ is absorbed and some is reflected diffusely causing multiple reflections to occur. This makes radiation between grey surfaces somewhat complex compared with black surfaces since, is the rate at which radiation leaves per unit area of surface i over the whole span of wavelengths. If the incident radiation arriving at a grey surface i in an enclosure is \mathcal{Q}_{li} , termed radiosity and the total radiosity Q_{0i} , which in the SI system has the units W/m², corresponding to a flux $q_{li} = Q_{li}/A_i$, then the reflected flux, that is, energy per unit area, is $\mathbf{r}_i q_{Ii}$. The emitted flux is $\mathbf{e}_i E_{bi} = \mathbf{e}_i \sigma T_i^4$ and the radiosity is then given by:

$$q_{0i} = \mathbf{e}_i E_{bi} + \mathbf{r}_i q_{Ii} \tag{9.1}$$

The net radiation from the surface is given by:

$$Q_i$$
 = (rate at which energy leaves the surface)
- (rate at which energy arrives at the surface)

ë.

$$Q_i = q_{0i} - Q_{1i} = A_i(q_{0i} - q_{1i})$$
 (9.139)

Substituting from equation 9.138 in equation 9.139 and noting that $(e_i + r_i) = 1$, then:

$$Q_i = A_i e_i E_{bi} / r_i + (A_i / r_i) [q_{0i} (1 - e_i) - q_{0i}]$$

$$= (A_i e_i / r_i) (E_{bi} - q_{0i})$$
(9.140)

If the temperature of a grey surface is known, then the net heat transfer to or from the surface may be determined from the value of the radiosity qo. With regard to signs, the usual convention is that a positive value of Q_i indicates heat transfer from grey surfaces.

Example 9.20

HEAT TRANSFER

Radiation arrives at a grey surface of emissivity 0.75 at a constant temperature of 400 K, at the rate of 3 kW/m². What is the radiosity and the net rate of radiation transfer to the surface? What coefficient of heat transfer is required to maintain the surface temperature at 300 K if the rear of the surface is perfectly insulated and the front surface is cooled by convective heat transfer to air at 295 K?

Solution

Since
$$e + r = 1$$
: $r = 0.25$. . . From equation 9.118: $E_b = (5.67 \times 10^{-8} \times 400^4) = 1452 \text{ W/m}^2$ From equation 9.138: $q_0 = eE_b + rq_I$

=
$$(0.75 \times 1452) + (0.25 \times 3000) = 1839 \text{ W/m}^2$$

From equation 9.140:
$$Q/A = q = (1.0 \times 0.75/0.25)(1452 - 1839) = -1161 \text{ W/m}^2$$

where the negative value indicates heat transfer to the surface.

For convective heat transfer from the surface:

 $q_c = h(T_s - T_{ambient})$

$$h_c = q_c/(T_s - T_{\text{ambient}}) = 1161/(400 - 295) = 11.1 \text{ W/m}^2 \text{ K}$$

and:

radiation with each other only, then, assuming $T_1 > T_2$, Q_{12} is the net rate of transfer For the simplest case of a two-surface enclosure in which surfaces 1 and 2 exchange from 1, Q_1 or the rate of transfer to 2, Q_2 .

Thus:

$$Q_{12} = Q_1 = -Q_2 (9.141)$$

Substituting from equation 9.139:

$$Q_1 = A_1(q_{01} - q_{11}) (9.142)$$

$$q_{11}A_1 = q_{01}A_1F_{11} + q_{02}A_2F_{21}$$
 (9.143)

that is:

and:

(rate of energy incident upon surface 1)

= (rate of energy arriving at surface 1 from itself)

+ (rate of energy arriving at surface 1 from surface 2)

From equations 9.142 and 9.143 and using $A_1F_{12} = A_2F_{21}$, then:

$$Q_1 = q_{01}(A_1 - A_1 F_{11}) - q_{02} A_1 F_{12}$$
 (9.144)

Since, by the summation rule, $(A_1 - A_1F_{11}) = A_1F_{12}$, then:

$$Q_1 = (A_1 F_{12})(q_{01} - q_{02}) \tag{9.145}$$

From equation 9.140:

$$Q_1 = (A_1 e_1/r_1)(E_{b1} - q_{01})$$
 and $Q_2 = (A_2 e_2/r_2)(E_{b2} - q_{02})$ (9.146)

Substituting from equation 9.146 into equation 9.145 and using the relationships in equation 9.141 gives:

$$Q_{12}[(1/A_1F_{12}) + (\mathbf{r}_1/A_1\mathbf{e}_1) + (\mathbf{r}_2/A_2\mathbf{e}_2)] = (E_{b1} - E_{b2})$$

and hence
$$Q_{12} = (E_{b1} - E_{b2})/[(1/A_1F_{12}) + (\mathbf{r}_1/A_1\mathbf{e}_1) + (\mathbf{r}_2/A_2\mathbf{e}_2)]$$
 (9.147)

and:

Since $\mathbf{r} = 1 - \mathbf{e}$, then, writing $E_{b1} = \sigma T_1^4$ and $E_{b2} = \sigma T_2^4$.

$$Q_{12} = [\sigma(T_1^4 - T_2^4)]/[(1/A_1F_{12}) + (1 - e_1)/(A_1e_1) + (1 - e_2)/(A_2e_2)]$$
(9.148)

Equation 9.148 is the same as equation 9.127 for black body exchange with two additional terms $(1 - e_1)/(A_1e_1)$ and $(1 - e_2)/(A_2e_2)$ introduced in the denominator for surfaces 1 and 2.

Radiation between parallel plates

For two large parallel plates of equal areas, and separated by a small distance, it may be assumed that all of the radiation leaving plate 1 falls on plate 2, and similarly all of the radiation leaving plate 2 falls on plate 1.

$$F_{12} = F_{21} = 1$$

Thus:

and:

 $A_1 = A_2$

Substituting in equation 9.148:

$$Q_{12} = \frac{A_1 \sigma (T_1^4 - T_2^4)}{1 + (1 - \epsilon_1)/\epsilon_1 + (1 - \epsilon_2)/\epsilon_2}$$

 $\frac{Q_{12}}{A_1} = q_{12} = \frac{\sigma(T_1^4 - T_2^4)}{1 + (1 - e_2)/e_2}$

and:

(9.149)

Other cases of interest include radiation between:

- (i) two concentric spheres
- (ii) two concentric cylinders where the length: diameter ratio is large.

In both of these cases, the inner surface 1 is convex, and all the radiation emitted by it falls on the outer surface 2.

Thus:

 $F_{12} = 1$

and from the reciprocal rule:

$$F_{21} = F_{12} \frac{A_1}{A_2} = \frac{A_1}{A_2}$$
 (equation 9.126)

Substituting in equation 9.148:

$$Q_{12} = \frac{A_1 \sigma(T_1^4 - T_2^4)}{1 + [(1 - \mathbf{e}_1)/\mathbf{e}_1] + [(1 - \mathbf{e}_2)/\mathbf{e}_2] \frac{A_1}{A_2}}$$

HEAT TRANSFER

$$= \frac{A_1 \sigma(T_1^4 - T_2^4)}{\frac{1}{\mathbf{e}_1} + \frac{1}{\mathbf{e}_2} (1 - \mathbf{e}_2) \frac{A_1}{A_2}}$$

$$\frac{Q_{12}}{A_1} = q_{12} = \frac{\sigma(T_1^4 - T_2^4)}{\frac{(1 - \mathbf{e}_2) A_1}{\mathbf{e}_1}}$$
(9.150)

For radiation from surface 1 to extensive surroundings $(A_1/A_2 \rightarrow 0)$, then:

$$q_{12} = \mathbf{e}_1 \sigma(T_1^4 - T_2^4)$$
 (9.151)

Radiation shield

The rate of heat transfer by radiation between two surfaces may be reduced by inserting a shield, so that radiation from surface 1 does not fall directly on surface 2, but instead is intercepted by the shield at a temperature T_{sh} (where $T_1 > T_{sh} > T_2$) which then reradiates to surface 2. An important application of this principle is in a furnace where it is necessary to protect the walls from high-temperature radiation.

The principle of the radiation shield may be illustrated by considering the simple geometric configuration in which surfaces 1 and 2 and the shield may be represented by large planes separated by a small distance as shown in Figure 9.43.

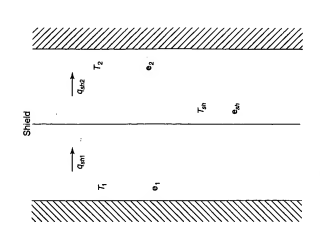


Figure 9.43. Radiation shield

Neglecting any temperature drop across the shield (which has a surface emissivity e_{3h}), then in the steady state, the transfer rate of radiant heat to the shield from the surface I must equal the rate at which heat is radiated from the shield to surface 2. Application of equation 9.149 then gives:

$$q_{sh} = q_{sh1}(=q_{sh2}) = \frac{\sigma(T_1^4 - T_{sh}^4)}{\binom{1}{e_1} + \binom{1}{\binom{1}{e_{sh}}} - 1} = \frac{\sigma(T_{sh}^4 - T_2^4)}{\binom{1}{e_{sh}} + \binom{1}{\binom{1}{e_2}} - 1}$$
(9.152)

Eliminating T_{sh}^4 in terms of T_1^4 and T_2^4 from equation 9.152:

$$T_1^4 - T_{sh}^4 = \frac{q_{sh}}{\sigma} \left(\frac{1}{\mathbf{e}_1} + \frac{1}{\epsilon_{sh}} - 1 \right)$$

$$T_{sh}^4 - T_2^4 = \frac{q_{sh}}{\sigma} \left(\frac{1}{\mathbf{e}_{sh}} + \frac{1}{\epsilon_2} - 1 \right)$$

$$T_1^4 - T_2^4 = \frac{q_{sh}}{\sigma} \left(\frac{1}{\mathbf{e}_{sh}} + \frac{1}{\epsilon_2} - 1 \right)$$

 $T_1^4 - T_2^4 = \frac{q_{sh}}{\sigma} \left(\frac{1}{\mathbf{e}_1} + \frac{2}{\mathbf{e}_{sh}} + \frac{1}{\mathbf{e}_2} - 2 \right)$

Adding:

and:

(9.153)

Then from equations 9.149 and 9.153;

$$\frac{q_{sh}}{q_{12}} = \frac{\left(\frac{1}{e_1}\right) + \left(\frac{1}{e_2}\right) - 1}{\frac{1}{e_1} + \frac{1}{e_2} + \frac{1}{e_3} - 2}$$
(9.154)

For the special case where all the emissivities are equal $(\mathbf{e}_1 = \mathbf{e}_{st} = \mathbf{e}_2)$:

$$\frac{q_{sh}}{q_{12}} = 1/2$$

Similarly, it can be shown that if n shields are arranged in series, then:

$$\frac{q_{sh}}{q_{12}} = \frac{1}{n+1}$$

surroundings will have been reduced to q_{sh} . The extent to which T_2 is reduced depends In practice, as a result of introducing the radiation shield, the temperature T_2 will fall because a heat balance must hold for surface 2, and the heat transfer rate from it to the on the heat transfer coefficient between surface 2 and the surroundings.

Multi-sided enclosures

For the more complex case of a multi-sided enclosure formed from n surfaces, the radiosities may be obtained from an energy balance for each surface in turn in the enclosure. Thus the energy falling on a typical surface i in an enclosure formed from

HEAT TRANSFER

n surfaces is:

$$A_iq_{1i} = q_{01}A_1F_{1i} + q_{02}A_2F_{2i} + q_{03}A_3F_{3i} + \dots + q_{0n}A_nF_{ni}$$
 (9.155)

which is intercepted by surface i. $q_{01}A_1 = q_{01}$ is the energy leaving surface 1 and F_{1i} is where $A_i q_{Ii} = Q_i$ is the energy incident upon surface i, $q_{01} A_1 F_{1i}$ is the energy leaving surface 1 which is intercepted by surface i and $q_{0n}A_nF_{ni}$ is the energy leaving surface n the fraction of this which is intercepted by surface i.

From equation 9.138:

$$q_{Ii} = (q_{Oi} - \mathbf{e}_i E_{bi})/\mathbf{r}$$

Substituting for q_{II} into equation 9.155 gives:

$$A_i(q_{0i} - \mathbf{e}_i E_{bi})/\mathbf{r}_i = A_1 F_{1i} q_{01} + A_2 F_{2i} q_{02} + A_3 F_{3i} q_{03}$$

$$+ \dots + A_j F_{ji} q_{0i} + \dots + A_n F_{ni} q_{0n}$$
 (9.156)

Noting, for example, that for surface 2, i = 2, then:

$$A_2(q_{02} - \mathbf{e}_2 E_{b2})/\mathbf{r}_2 = A_1 F_{12} q_{01} + A_2 F_{22} q_{02} + A_3 F_{32} q_{03}$$

$$+ \cdots + A_j F_{j2} q_{0j} + \cdots + A_n F_{n2} q_{on}$$
 (9.157)

Rearranging

$$A_1 F_{12} q_{01} + [A_2 F_{22} - (A_2/r_2)] q_{02} + A_3 F_{32} q_{03} + \dots + A_j F_{j2} q_{0i} + \dots + A_n F_{n2} q_{0n} = (A_2 e_2/r_2) E_{b2}$$

$$(9.158)$$

then be solved for the unknown radiosities, $q_{01}, q_{02}, \dots q_{0n}$. The radiation heat transfer is then obtained from equation 9.140. This approach requires data on the areas and view Equations similar to equation 9.158 may be obtained for each of the surfaces in an factors for all pairs of surfaces in the enclosure and the emissivity, reflectivity and the black body emissive power for each surface. Should any surface be well insulated, then, enclosure, i = 1, i = 2, i = 3, i = n and the resulting set of simultaneous equations may in this case, $Q_i = 0$ and:

$$A_i(\mathbf{e}_i/\mathbf{r}_i)(E_{bi}-q_{Oi})=0$$

Since, in general, $A_i(\mathbf{e}_i/\mathbf{r}_i) \neq 0$, then $E_{bi} = q_{Oi}$.

If a surface has a specified net thermal input flux, say q11, then, from equation 9.140.

$$E_{bi} = (\mathbf{r}_i/(A_i\mathbf{e}_i))q_{Ii} + q_{Oi}.$$

It may be noted that this approach assumes that the surfaces are grey and diffuse, that emissivity and reflectivity do not vary across a surface and that the temperature, irradiation and radiosity are constant over a surface. Since the technique uses average values over noting that a number of surfaces may be regarded as a single surface, that it may be necessary to split one surface up into a number of smaller surfaces and also possibly to introduce an imaginary surface into the system, to represent the surroundings, for example. a surface, the subdivision of the enclosure into surfaces must be undertaken with care, In a real situation, there may be both grey and black surfaces present and, for the latter, \mathbf{r}_i tends to zero and (A_i/\mathbf{r}_i) and $(A_i\mathbf{e}_i/\mathbf{r}_i)$ become very large.

Example 9.21

A horizontal circular plate, 1.0 m in diameter, is to be maintained at 500 K by placing it 0.20 m directly beneath a horizontal electrically heated plate, also 1.0 m in diameter, maintained at 1000 K. The assembly is exposed to black surroundings at 300 K, and convection heat transfer is negligible. Estimate the electrical input to the heater and the net rate of heat transfer to the plate if the emissivity of the heater is 0.75 and the emissivity of the plate 0.5.

Solution

Taking surface 1 as the heater, surface 2 as the heated plate and surface 3 as an imaginary enclosure consisting of a vertical cylindrical surface representing the surroundings, then, for each surface:

Surface	A (m ²)	Ð	-	(A/r)	(Ae/r) (m ²)
3 2 -	1.07 1.07 0.628	0.75 0.50 1.0	0.25	4.28	3.21

For surface 1:

For a plane surface: $F_{11} = 0$ and $A_1F_{11} = 0$

Using the nomenclature of Figure 9.39: For co-axial parallel discs with $r_1=r_2=0.5~{\rm m}$ and $L=0.2~{\rm m}$:

 $R_1 = r_1/L = (0.5/0.20) = 2.5$

 $R_2 = r_2/L = (0.5/0.20) = 2.5$

 $S = 1 + (1 + R_2^2)/R_1^2 = 1 + (1 + 2.5^2)/2.5^2 = 2.16$

 $F_{12} = 0.5\{S - [S^2 - 4(r_2/r_1)^2]^{0.5}\}$

From Figure 9.39ii:

and:

 $= 0.5 \times \{2.16 - [(2.16^2 - 4(0.5/0.5)^2]^{0.5}\} = 0.672$

 $A_1F_{12} = (1.07 \times 0.672) = 0.719 \text{ m}^2$

and, from the summation rule:

 $A_1F_{13} = A_1 - (A_1F_{11} + A_1F_{12}) = 1.07 - (0 + 0.719) = 0.350 \text{ m}^2$

For surface 2:

For a plane surface:

 $A_2F_{22}=0$

and by the reciprocity rule:

 $A_2F_{21} = A_1F_{12} = 0.719 \text{ m}^2$

 $A_2F_{23} = A_1F_{13} = 0.350 \text{ m}^2$

 $A_3F_{31} = A_1F_{13} = 0.350 \text{ m}^2$

 $A_3F_{32} = A_2F_{23} = 0.350 \text{ m}^2$

From the summation rule:

From equation 9.112:

By the reciprocity rule:

For surface 3:

By symmetry:

 $= 0.785 - (0.350 + 0.350) = 0.085 \text{ m}^2$

 $A_3F_{33} = A_3 - (A_3F_{31} + A_3F_{32})$

 $= 5.67 \times 10^4 \text{ W/m}^2 \text{ or } 56.7 \text{ kW/m}^2$ $E_{b1} = \sigma T_1^4 = (5.67 \times 10^{-8} \times 1000^4)$

HEAT TRANSFER

$$E_{b2} = \sigma T_2^4 = (5.67 \times 10^{-8} \times 500^4)$$

 $= 3.54 \times 10^3 \text{ W/m}^2 \text{ or } 3.54 \text{ kW/m}^2$ $E_{b3} = \sigma T_3^4 = (5.67 \times 10^{-8} \times 300^4)$ $= 0.459 \times 10^3 \text{ W/m}^2 \text{ or } 0.459 \text{ kW/m}^2$

 $q_{03} = E_{b3} = 0.459 \text{ kW/m}^2$

Since surface 3 is a black body,

From equations 9.157 and 9.158:

 $(A_1F_{11} - A_1/r_1)q_{01} + A_2F_{21}q_{02} + A_3F_{31}q_{03} = -E_{b1}A_1e_1/r_1$

 $\times (0 - 4.28)q_{01} + 0.719q_{02} + (0.350 \times 0.459)$ $= -(56.7 \times 1.07 \times 0.75)/0.25$

 $0.719q_{02} - 4.28q_{01} = -182$

Ξ

3

 $(A_1F_{12}q_{01}) + (A_2F_{22} - A_2/r_2)q_{02} = -E_{b2}A_2e_2/r_2$

and: 9

 $0.719q_{01} + (0 - 1.07/0.5)q_{02} = -(3.54 \times 1.07 \times 0.5)/0.5$

 $0.719q_{O1} - 2.14q_{O2} = -3.79$

Solving equations 1 and 2 simultaneously gives:

ä

 $q_{01} = 45.42 \text{ kW/m}^2$ and $q_{02} = 17.16 \text{ kW/m}^2$

power input to the heater = rate of heat transfer from the heater

From equation 9.140:

 $Q_1 = (A_1 \mathbf{e}_1/\mathbf{r}_1)(E_{b1} - q_{O1}) = (1.07 \times 0.75/0.25)(56.7 - 45.42) = 36.2 \text{ kW}$

Again, from equation 9.140, the rate of heat transfer to the plate is:

 $Q_2 = (A_2e_2/r_2)(E_{b2} - q_{02}) = (1.07 \times 0.5/0.25)(3.54 - 17.16) = -14.57 \text{ kW}$

where the negative sign indicates heat transfer to the plate.

9.5.6. Radiation from gases

In the previous discusion surfaces have been considered which are isothermal, opaque and grey which emit and reflect diffusely and are characterised by uniform surface radiosity in that it neither absorbs nor scatters the surface radiation nor does it emit radiation itself. Whilst, in most cases, such assumptions are valid and permit reasonably accurate results to be calculated, there are occasions where such assumptions do not hold and more refined techniques are required such as those described in the specialist literature (45,48-53). For non-polar gases such as N2 and O2, the foregoing assumptions are largely valid, since the gases do not emit radiation and they are essentially transparent H₂O vapour, NH₃ and hydrocarbon gases, however, since these not only emit and absorb over a wide temperature range, but also radiate in specific wavelength intervals and the medium separating the surfaces has been assumed to be non-participating, to incident radiation. This is not the case with polar molecules such as CO2 and called bands. Furthermore, gaseous radiation is a volumetric rather than a surface phenonemon. HEAT TRANSFER

Whilst the calculation of the radiant heat flux from a gas to an adjoining sufface developed by HOTTEL and MANGLESDORF⁽⁵⁴⁾, which involves the determination of to a surface element, d41, near the centre of the base of the hemisphere. Emission from embraces inherent spectral and directional effects, a simplified approach has been radiation emission from a hemispherical mass of gas of radius L. at temperature he gas per unit area of the surface is then:

$$E_g = e_g \sigma T_g^* \tag{9.159}$$

where the gas emissivity e_s is a function of T_s , the total pressure of the gas P, the partial

pressure of the radiating gas P_g and the radius of the hemisphere L.

9.45. Although these data refer to water vapour or carbon dioxide alone in a mixture of also given in the figure must be used. Similar data for carbon dioxide are given in Figure Data on the emissivity of water vapour at a total pressure of 101.3 kN/m2 are plotted non-radiating gases, they may be extended to situations where both are present in such a in Figure 9.44 for different values of the product of the vapour partial pressure P_{w} and he hemisphere radius L. For other values of the total pressure, the correction factor C. mixture by expressing the total emissivity as:

$$\mathbf{e}_{g} = \mathbf{e}_{w} + \mathbf{e}_{c} - \Delta \mathbf{e} \tag{9.160}$$

where Δe is a correction factor, shown in Figure 9.46, which allows for the reduction in

by using the concept of mean beam length L, which correlates the dependence of gas Essentially the mean beam length is the radius of the hemisphere of gas whose emissivity is equivalent to that in the particular geometry considered, and typical values of L, which are then used to replace L in Figures 9.38-40 are shown in Table 9.7. Using these data and Figures 9.38-9.40, the rate of transfer of radiant heat to a surface of area A, due to radiating to an element at the centre of the base, they may be extended to other geometries emissivity with both the size and shape of the gas geometry in terms of a single parameter. Although these data provide the emissivity of a hemispherical gas mass of radius emission associated with mutual absorption of radiation between the two species. emission from an adjoining gas is given by:

$$Q = \mathbf{e}_{\kappa} A_{s} \sigma T_{\kappa}^{*} \tag{9.161}$$

A black surface will not only absorb all of this radiation but will also emit radiation, and the net rate at which radiation is exchanged between the gas and the surface at temperature T, is given by:

$$Q_{net} = A_s \sigma(\mathbf{e}_g T_g^s - \mathbf{a}_g T_g^s) \tag{9.162}$$

in this equation, the absorptivity ag may be obtained from the emissivity using expressions of the form(54);

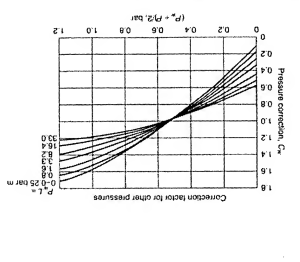
$$\mathbf{a}_{w} = C_{w} \mathbf{e}_{w} (T_{g} / T_{s})^{0.43}$$
 (9.163)

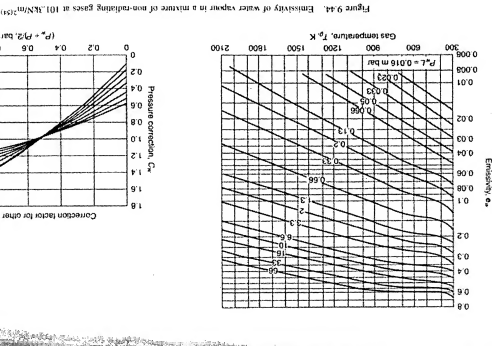
water

$$\mathbf{a}_c = C_c \mathbf{e}_c (T_g/T_s)^{0.65}$$

carbon dioxide:
$$\mathbf{a}_c = C_c \mathbf{e}_c (T_g/T_s)^{0.65}$$
 (9.164)

where e., and C., and e. and C. are obtained from Figures 9.38 and 9.39 respectively noting that T_g is replaced by T_s and $(P_w L_r)$ or $(P_c L_s)$ by $[P_w L_r (T_s/T_g)]$ or $[P_c L_c (T_s/T_g)]$





469

HEAT TRANSFER

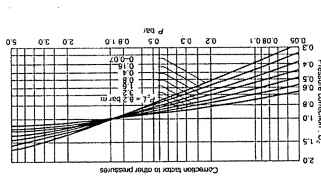


Figure 9.45. Emissivity of carbon dioxide in a mixture of non-radiating gases at 101.3 kN/mx^{2,640} 900 1200 1500 Gas temperature, T_R K

\$100

1800

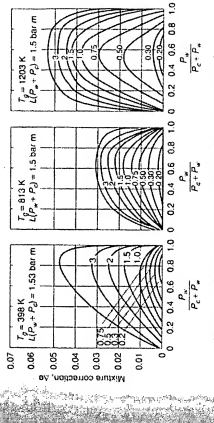


Figure 9.46. Currection factor for water vapour-carbon diaxide mixtures (54)

respectively. It may be noted that, in the presence of both water vapour and carbon dioxide, the total absorptivity is given by:

$$\mathbf{a}_g = \mathbf{a}_w + \mathbf{a}_c - \Delta \mathbf{a} \tag{9.1}$$

where $\Delta u = \Delta e$ is obtained from Figure 9.46. If the surrounding surface is grey, some of the radiation may be reflected and equation 9.162 may be modified by a factor, $e_i/(1-(1-a_g)(1-e_g))$ to take this into account. This leads to the following equation for the heat transferred per unit time from the gas to the surface:

$$Q = \sigma \mathbf{e}_s A_s (\mathbf{e}_g T_g^4 - \mathbf{a}_g T_s^4) / [1 - (1 - \mathbf{a}_g)(1 - \mathbf{e}_g)]$$
 (9.166)

Table 9.7. Mean beam lengths for various geometries (54)

Geometry	Characteristic length	Mean beam length, L,
Sphere — radiation to surface Infinite circular cylinder —	Diameter, D	0.650
radiation to curved surface Semi-infinite evlinder—	Diameter, D	0.950
radiation to base Cylinder of equal height and	Diameter, D	0.650
diameter — radiation to emire surface Infinite parallel planes —	Diameter, D	0.600
radiation to planes	Spacing between planes, L.	1.807
Cube — radiation to any surface Shape of volume, V —	Side, L	799.0
radiation to surface of area, A	Ratio; volume/area, (V/A)	3.6(V/A)

____1 100.0 00E

O.002 800.0 \$00.0 \$00.0

10,0 800.0

£0.0

1000 90.0 80.0 1.0

S.0

Emissivity, e, 0.02

009

1

Example 9.22

The walls of a combustion chamber, 0.5 m in diameter and 2 m long, have an emissivity of 0.5 and are maintained at 750 K. If the combustion products containing 10 per cent carbon dioxide and 10 per cent water-vapour are at 150 kN/m² and 1250 K, what is the net rate of radiation to the walls?

Solution

The partial pressures of carbon dioxide (Pe) and of water (Pw) are:

$$P_c = P_w = (10/100)150 = 15.0 \text{ kN/m}^2 \text{ or } (15.0/100) = 0.15 \text{ bar}$$

From Table 9.7:

$$L_{\rm r} = 3.6 V/\Lambda = 3.6 (\pi/4 \times 0.5^2 \times 2)/(2\pi/4 \times 0.5^2) + (0.5\pi \times 2.0) = 0.4 \text{ m}$$

For water vapour:

$$P_{\rm s}L_{\rm s} = (0.15 \times 0.4) = 0.06$$
 bar m

and from Figure 9.44, e., = 0.075

$$P = (150/100) = 1.5 \text{ bar. } P_w = 0.15 \text{ bar and: } 0.5(P_w + P) = 0.825 \text{ bar}$$

Since $P_{\mu}L_{\mu} = 0.06$ bar m, then from Figure 9.44;

$$C_{\rm s} = 1.4$$
 and $\epsilon_{\rm s} = (1.4 \times 0.075) = 0.105$

Fur curbon dioxide:

$$P_c I_r = (0.15 \times 0.4) = 0.06$$
 bar m

and from Figure 9.45, e. = 0.037 Since P=1.5 bar, $P_c=0.15$ bar and $P_cL_{\varphi}=0.06$ bar in, then, from Figure 9.38:

$$C_c = 1.2$$
 and $e_r = (1.2 \times 0.037) = 0.044$

$$(P_w + P_c)I_c = (0.15 + 0.15)0.4 = 0.12$$
 but

$$P_c/(P_c + P_w) = 0.15/(0.15 + 0.15) = 0.5$$

Thus, from Figure 9.45 for $T_g > 1203$ K. $\Delta e = 0.001$

and, from equation 9.160;

 $c_g = c_w - \Delta c = (0.105 + 0.044 - 0.001) = 0.148$

For water vapour:

$$P_w L_v(T_s/T_g) = 0.06(750/1250) = 0.036$$
 bar m

and, from Figure 9.44 at 750 K, $\epsilon_{\rm w}=0.12$ Since $0.5(P_w+P)=0.825$ bar and $P_w L_d(T_s/T_g)=P_c L_{\rm w}(T_s/T_g)=0.036$ bar m, then, from Figure 9.44; $C_w=1.40$ and $\epsilon_{\rm w}=(0.12\times1.40)=0.168$ and the absorptivity, from equation 9.163 is:

$$\mathbf{a}_{s} = \mathbf{e}_{s}(T_{s}/T_{s})^{0.45} = 0.168(1250/750)^{0.45} = 0.212$$

For curbon dioxide:

From Figure 9.45 at 750 K, $\epsilon_r=0.08$ From Figure 9.45 at P=1.5 bar and $P_cL_c(T_s/T_g)=0.036$ bar m:

 $C_c = 1.02$ and $e_c = (0.08 \times 1.02) = 0.082$

and the absorptivity, from equation 9.164 is:

$$\mathbf{a}_{v} = \mathbf{e}_{c}(T_{g}/T_{v})^{0.65} = 0.082(1250/750)^{0.65} = 0.114$$

$$P_{w}/(P_{c} + P_{w}) = 0.5$$
 and $(P_{c} + P_{w})I_{w}(T_{s}/T_{g}) = (0.036 + 0.036) = 0.072$ bur m

This, from Figure 9.46, for $T_r = 813$ K. $\Delta e = \Delta a < 0.01$ and this may be neglected.

$$a_t = a_w + a_t - \Delta a = (0.212 + 0.114 - 0) = 0.326$$

If the surrounding surface is black, then:

Ě

$$Q = aA_1(c_sT_s^4 - a_kT_s^4)$$

= $(5.67 \times 10^{-8} [(2(\pi/4)0.5^2) + (0.5\pi \times 2.0)])[(0.148 \times 1250^4) - (0.326 \times 750^4)]$

(equation 9.162)

$$= 5.03 \times 10^4 \text{ W} = 50.3 \text{ kW}$$

For grey walls, the correction factor allowing for multiple reflection of incident radiation is:

$$C_k = e_k/[1 - (1 - a_k)(1 - e_k)] = 0.5/[1 - (1 - 0.326)(1 - 0.5)] = 0.754$$

and hence: net radiation to the walls, $Q_{\kappa} = (50.3 \times 0.754) = 37.9 \text{ kW}$

Radiation from gases containing suspended particles

The estimation of the radiation from pulverised-fuel flames, from dust particles in flames and from flames made luminous as a result of the thermal decomposition of hydrocarbons io soot, involves an evaluation of radiation from clouds of particles. In pulverised-fuel flames, the mean particle size is typically 25 µm and the composition varies from a very high carbon content to virtually pure ash. In contrast, the suspended matter in luminous flames, resulting from soot formation due to incomplete mixing of hydrocarbons with air before being heated, consists of carbon together with very heavy hydrocarbons with an initial particle size of some $0.3 \mu m$. In general, pulverised-fuel particles are sufficiently large to be substantially opaque to incident radiation, whilst the particles in a luminous flame are so small that they act as semi-transparent bodies with respect to thermal or long wavelength radiation.

According to SCHACK⁽³⁵⁾, a single particle of soot transmits approximately 95 per cent of the incident radiation and a cloud must contain a very large number of particles before an appreciable emission can occur. If the concentration of particles is K', then the product of K' and the thickness of the layer L is equivalent to the product P_gL_r in the radiation of gases. For a known or measured emissivity of the flame e_f , the heat transfer rate per unit time to a wall is given by:

$$Q = e_f e_s \sigma(T_f^4 - T_w^4)$$
 (9.167)

where e_s is the effective emissivity of the wall, and T_f and T_w are the temperatures of the flame and wall respectively, e_f varies, not only from point to point in a flame, but also depends on the type of fuel, the shape of the burner and combustion chamber, and on the air supply and the degree of preheating of the air and fuel.

9.6. HEAT TRANSFER IN THE CONDENSATION OF VAPOURS

9.6.1. Film coefficients for vertical and inclined surfaces

When a saturated vapour is brought into contact with a cool surface, heat is transferred from the vapour to the surface and a film of condensate is produced. In considering the heat that is transferred, the method first put forward by NUSSELT $^{(56)}_2^{\circ}$ and later modified by subsequent workers is followed. If the vapour condenses on \mathbf{a}^4 although it is retarded by the viscosity of the liquid. The flow will normally be streamline and the heat flows through the film by conduction. In Nusselt's work it is assumed that the temperature of the film at the cool surface is equal to that of the surface, and at the vertical surface, the condensate film flows downwards under the influence of gravity, other side was at the temperature of the vapour. In practice, there must be some small difference in temperature between the vapour and the film, although this may generally be neglected except where non-condensable gas is present in the vapour.

It is shown in Chapter 3, that the mean velocity of a fluid flowing down a surface inclined at an angle ϕ to the horizontal is given by:

$$u = \frac{\rho g \sin \phi s^2}{3\mu}$$
 (equation 3.87)

 $\sin \phi = 1 \quad \text{and} \quad u = \frac{\rho g s^2}{3\mu}$ For a vertical surface:

The maximum velocity u_s which occurs at the free surface is:

$$u_s = \frac{\rho g \sin \phi \, s^2}{2\mu} \tag{equation 3.88}$$

and this is 1.5 times the mean velocity of the liquid.

difference in the mass rates of flow at distances x and x + dx from the top of the surface Since the liquid is produced by condensation, the thickness of the film will be zero at the top and will gradually increase towards the bottom. Under stable conditions the will result from condensation over the small element of the surface of length dx and width ω , as shown in Figure 9.47

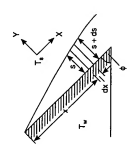


Figure 9.47. Condensation on an inclined surface

If the thickness of the liquid film increases from s to s + ds in that distance, the increase rate of flow of liquid dG is given by:

$$= \frac{d}{ds} \left(\frac{\rho^2 g \sin \phi s^3 w}{3\mu} \right) ds$$
$$= \frac{\rho^2 g \sin \phi}{\mu} ws^2 ds$$

If the vapour temperature is T_s and the wall temperature is T_w , the heat transferred by hermal conduction to an element of surface of length dx is:

$$\frac{k(T_s - T_w)}{\varepsilon} w \, \mathrm{d}x$$

where k is the thermal conductivity of the condensate. Thus the mass rate of condensation on this small area of surface is:

$$\frac{k(T_s - T_w)}{c_1} w dx$$

where λ is the latent heat of vaporisation of the liquid.

$$\frac{k(T_s - T_w)}{s\lambda} w \, dx = \frac{\rho^2 g \sin \phi}{\mu} w s^2 \, ds$$

On integration:

$$\mu k(T_s - T_w)x = \frac{1}{4}\rho^2 g \sin \phi \, s^4 \lambda$$

since s = 0 when x = 0.

$$s = \sqrt{\frac{4\mu kx(T_s - T_w)}{g\sin\phi \lambda \rho^2}}$$
 (9.168)

by the heat transfer coefficient h at x = x, = k/s, and hence:

$$h = \sqrt{\frac{\rho^2 g \sin \phi \, \lambda k^3}{4 \mu x (T_s - T_w)}} \tag{9.169}$$

$$Nu = \frac{hx}{k} = \sqrt{\frac{\rho^2 g \sin \phi \lambda x^3}{4 \mu k (T_s - T_w)}}$$
 (9.170)

Traises from a theoretical value of infinity at the top as the condensate film thickens. Since an value of the heat transfer coefficient over the whole surface, between x=0Expressions give point values of h and Nu_x at x=x. It is seen that the coefficient

$$h_m = \frac{1}{x} \int_0^x h \, dx = \frac{1}{x} \int_0^x Kx^{-1/4} \, dx \quad \text{(where } K \text{ is independent of } x\text{)}$$

$$\vdots$$

$$= \frac{1}{x} \frac{x^{3/4}}{3} = \frac{4}{3} Kx^{-1/4} = \frac{4}{3}h$$

$$= 0.943 \sqrt{\frac{\rho^2 g \sin \phi \lambda k^3}{\mu x \Delta T_f}}$$

(9.171)

 $\Delta T_{\rm e}$ is the temperature difference across the condensate film. A vertical surface, $\sin \phi = 1$ and:

ertical surface,
$$\sin \phi = 1$$
 and:

$$h_m = 0.943 \sqrt{\frac{\rho^2 g \lambda k^3}{\mu x \Delta T_f}}$$
 (9.172)

9.6.2. Condensation on vertical and horizontal tubes

The Nusselt equation

If vapour condenses on the outside of a vertical tube of diameter d_o , then the hydraulic mean diameter for the film is:

$$\frac{4 \times \text{flow area}}{\text{wetted perimeter}} = \frac{4S}{b} \quad \text{(say)}$$

If G is the mass rate of flow of condensate, the mass rate of flow per unit area G' is G/\hat{S}_{S}^{*} and the Reynolds number for the condensate film is then given by:

$$Re = \frac{(4S/b)(G/S)}{\mu} = \frac{4G}{\mu b} = \frac{4M}{\mu}$$
 (9.173)

where M is the mass rate of flow of condensate per unit length of perimeter, or:

$$M = \frac{G}{\pi d_o}$$

For streamline conditions in the film, $4M/\mu \neq 2100$ and:

$$h_m = \frac{Q}{A\Delta T_f} = \frac{G\lambda}{bI\Delta T_f} = \frac{\lambda M}{I\Delta T_f}$$

From equation 9.172:

$$h_m = 0.943 \left(\frac{k^3 \rho^2 g}{\mu} \frac{\lambda}{1 \Delta T_f} \right)^{1/4} = 0.943 \left(\frac{k^3 \rho^2 g}{\mu} \frac{h_m}{M} \right)^{1/4}$$

 $h_m \left(\frac{\mu^2}{k^3 \rho^2 g} \right)^{1/3} = 1.47 \left(\frac{4M}{\mu} \right)^{-1/3}$

(9.1

 $h_m = 0.72 \left(\frac{k^3 \rho^2 g \lambda}{d_o \mu \Delta T_f} \right)^{1/4}$ For horizontal tubes, Nusselt proposes the equation:

This may be rearranged to give:
$$h_m\left(\frac{\mu^2}{k^3\rho^2g}\right)^{1/3}=1.51\left(\frac{4M}{\mu}\right)^{-1/3}$$

where M is the mass rate of flow per unit length of tube.

that, provided the length is more than three times the diameter, the horizontal the tubes M = G/l, where l is the length of the tube. Comparison of the two equations equation for condensation, noting that for vertical tubes $M = G/\pi d_o$ and for holy This is approximately the same as equation 9.173 for vertical tubes and is a ju give a higher transfer coefficient for the same temperature conditions.

For j vertical rows of horizontal tubes, equation 9.175 may be modified to give

$$h_m = 0.72 \left(\frac{k^3 \rho^2 g \lambda}{j d_o \mu \Delta T_f} \right)^{1/4}$$

HEAT TRANSFER

FFRN(28) suggests that, based on the performance of commercial exchangers, this equation is too conservative and that the exponent of j should be nearer $-\frac{1}{5}$ than $-\frac{1}{4}$. his topic is discussed in Volume 6, Chapter 12.

Experimental values

the requirements of the theory. In particular, it is necessary for the condensate to form a uniform film on the tubes, for the drainage of this film to be by gravity, and the how streamline. Although some of these requirements have probably not been entirely surface, where precautions were taken to see that the conditions were met, are in very good agreement with Nusselt's theory. The results for most of the workers are within 15 per centifor horizontal tubes, although they tend to be substantially higher than the theoretical of yetical tubes. Typical values are given in Table 9.8 taken from MCADAMS⁽²⁷⁾ and Injecting Nusselt's equation it is important to ensure that the conditions comply with and so on, are sufficiently close to give support to the theory. Some data obtained by HASELDEN and PROSAD⁽⁵⁷⁾ for condensing oxygen and nitrogen vapours on a vertical fulfilled, results for pure vapours such as steam, benzene, toluene, diphenyl, ethanol elsewhere.

Table 9.8. Average values of film coefficients h_m for condensation of pure saturated vapours on horizontal tubes

	and dec		
Vapour	Value of h_m (W/m ² K)	Value of h_m (Btu/h ft² °F)	Range of ΔT_f (deg K)
Steam	10,000-28,000	1700-5000	1-11
Steam	18,000-37,000	3200-6500	4-37
Benzene	1400-2200	240-380	23-37
Diphenyl	1300-2300	. 220-400	4-15
Toluene	1100-1400	190-240	31-40
Methanol	2800-3400	200-600	8-16
Ethanol	1800-2600	320-450	6-22
Propanol	1400-1700	250-300	13-20
Oxygen	3300-8000	570-1400	0.08-2.5
Nitrogen	2300-5700	400-1000	0.15-3.5
Ammonia	0009	1000	1
Freon-12	1100-2200	200-400	ı

sidering commercial equipment, there are several factors which prevent the ins of Nusselt's theory being met. The temperature of the tube wall will not and for a vertical condenser with a ratio of ΔT at the bottom to ΔT at the he film coefficient should be increased by about 15 per cent.

of vapour velocity

ough the film may sometimes be disrupted mechanically as a result of the signall waves. For the downward flow of vapour, TEN BOSCH(58) has shown our velocity upwards tends to increase the thickness of the film and thus ises, considerably at high vapour velocities and may increase to two or three jie given by the Nusselt equation. It must be remembered that when a large 477

HEAT TRANSFER

fraction of the vapour is condensed, there may be a considerable change in velocity over the surface

When the vapour velocity is high, there will be an appreciable drag on the condensate Under conditions of high vapour velocity CARPENTER and COLBURN⁽⁵⁹⁾ have shown that turbulence may occur with low values of the Reynolds number, in the range 200-400 film and the expression obtained for the heat transfer coefficient is difficult to manage,

condensation at varying vapour velocities on the inner surface of a vertical tube which Carpenter and Colburn have put forward a simple correlation of their results

$$h_m = 0.065G'_m \sqrt{\frac{C_p \rho k(R'/\rho_{\nu} u^2)}{\mu \rho_{\nu}}}$$
(9.178)

where:

of the condensate and ho_v refers to the vapour. G_1' is the mass rate of flow per unit area at the top of the tube and G_2' the corresponding value at the bottom. R' is the shear stress. and u is the velocity calculated from G'_m . In this equation C_p , k, ρ , and μ are properties at the free surface of the condensate film.

for a number of organic vapours so that a plot of h_m and G'_m will provide a simple approximate correlation with separate lines for steam and for organic vapours as shown As pointed out by COLBURN⁽⁶⁰⁾, the group $C_{\rho}\rho k/\mu \rho_{v}$ does not vary very much in Figure 9.48^(60,61). Whilst this must be regarded as an empirical approximation it is very useful for obtaining a good indication of the effect of vapour velocity.

Turbulence in the film

h, will increase as a result of turbulence. The data of KIRKBRIDE (62) and BADGER (63,64) for the condensation of diphenyl vapour and Dowtherm on nickel tubes are expressed in If Re is greater than 2100 during condensation on a vertical tube the mean coefficient

$$h_m \left(\frac{\mu^2}{k^3 \rho^2 g} \right)^{1/3} = 0.0077 \left(\frac{4M}{\mu} \right)^{0.4}$$
 (9.179);

Comparing equation 9.176 for streamline flow of condensate and equation 9.179 for turbulent flow, it is seen that, with increasing Reynolds number, h_m decreases with streamline flow but increases with turbulent flow. These results are shown in Figure 9.49.

outside horizontal and vertical tubes, and the importance of avoiding flooding in vertical? Design equations are given in Volume 6, Chapter 12, for condensation both inside and tubes is stressed.

9.6.3. Dropwise condensation

In the discussion so far, it is assumed that the condensing vapour, on coming into contact with the cold surface, wets the tube so that a continuous film of condensate is formed. If the droplets initially formed do not wet the surface, after growing slightly

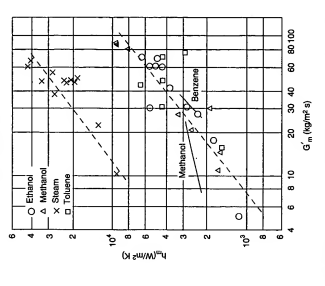


Figure 9.48. Average heat transfer data of CARPENTER and COLBURN⁽⁵⁹⁾ (shown as points) compared with those of TEPE and MUELLER⁽⁶¹⁾ (shown as solid lines). Dashed lines represent equation 9.178

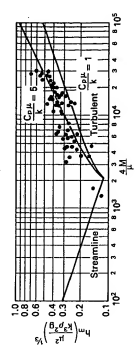


Figure 9.49. Effects of turbulence in condensate film

dropwise condensation has been obtained, and values of h from 40 to 114 kW/m² K have been obtained, with much higher values on occasions. This question has been discussed by DREW, NAGLE and SMITH(65) who have shown that there are many materials which make the surface non-wettable although, of these, only those which are firmly held to the they will fall from the tube exposing fresh condensing surface. This is known as dropwise condensation and, since the heat does not have to flow through a film by conduction, much higher transfer coefficients are obtained. Steam is the only pure vapour for which definite

surface are of any practical use. Mercaptans and oleic acid have been used to promote dropwise condensation, but at present there is little practical application of this technique coefficient, since for a condenser with steam, a value of about $11~\mathrm{kW/m^2~K}$ can be obtained Exceptionally high values of h will not give a corresponding increase in the overall with film condensation. On the other hand, it may be helpful in experimental work to reduce the thermal resistance on one side of a surface to a negligible value.

9.6.4. Condensation of mixed vapours

In the previous discussion it has been assumed that the vapour is a pure material, such as steam or organic vapour. If it contains a proportion of non-condensable gas and is cooled below its dew point, a layer of condensate is formed on the surface with a mixture of non-condensable gas and vapour above it. The heat flow from the vapour to the surface temperature difference. Secondly, since the concentration of vapour in the main stream is is then determined by the combination of these two effects, and its calculation requires a then takes place in two ways. Firstly, sensible heat is passed to the surface because of the greater than that in the gas film at the condensate surface, vapour molecules diffuse to the surface and condense there, giving up their latent heat. The actual rate of condensation knowledge of mass transfer by diffusion, as discussed in Chapter 10.

the method of COLBURN and HOUGEN(66) is considered. This requires a point-to-point calculation of the condensate-vapour interface conditions T_c and P_s . A trial and error In the design of a cooler-condenser for a mixture of vapour and a permanent gas, solution is required of the equation;

$$h_g(T_s - T_c) + \frac{q_v}{k_G \lambda(P_g - P_s)} = \frac{q_c}{h_o(T_c - T_{cm})} = U \Delta T$$
(9.180)

where the first term q_v represents the sensible heat transferred to the condensing surface, the second term q_{λ} the latent heat transferred by the diffusing vapour molecules, and, the third term qc the heat transferred from the condensing surface through the pipe wall, dirt and scales, and water film to the cooling medium, h_g is the heat transfer coefficient over the gas film, h_o the conductance of the combined condensate film, tube wall, dirt and scale films, and the cooling medium film and U the overall heat transfer coefficient. T_s is the vapour temperature, T_c the temperature of the condensate, T_{cm} the cooling medium temperature, ΔT the overall temperature difference = $(T_s - T_{cm})$, P_g is the partial pressure of diffusing vapour, P_s the vapour pressure at T_c , λ the latent heat of vaporisation per unit mass, and k_G the mass transfer coefficient in mass per unit time, unit area and unit partial pressure difference.

To evaluate the required condenser area, point values of the group $U\Delta T$ as a function of qc must be determined by a trial and error solution of equation 9.181. Integration of takes into account point variations in temperature difference, overall coefficient and mass a plot of q_c against $1/U\Delta T$ will then give the required condenser area. This method velocities and consequently produces a reasonably accurate value for the surface area

The individual terms in equation 9.181 are now examined to enable a trial solution to proceed. Values for h_g and k_G are most conveniently obtained from the CHILTON and

GOLBURN⁽⁶⁷⁾ analogy discussed in Chapter 10.

HEAT TRANSFER

$$h_g = \frac{j_h G' C_p}{(C_p \mu/k)^{0.67}} \tag{9.182}$$

$$k_G = \frac{j_d G'}{P_{Bm}(\mu/\rho D)^{0.67}} \tag{9.183}$$

Values of j_h and j_d are obtained from a knowledge of the Reynolds number at a given oint in the condenser. The combined conductance ho is evaluated by determining the $\mathbf{f}_{\text{ondensate}}$ film coefficient h_c from the Nusselt equation and combining this with the dirt ind tube wall conductances and a cooling medium film conductance predicted from the jeder-Tate relationships. Generally, ho may be considered to be constant throughout the exchanger.

From a knowledge of h_g , k_G , and h_o and for a given T_s and T_{cm} values of the condensate surface temperature T_c are estimated until equation 9.181 is satisfied. The calculations The repeated, and in this manner several point values of the group $U\Delta T$ throughout the condenser may be obtained.

complicated than the preceding single vapour-permanent gas case⁽⁶⁸⁾, and an example has been given by JEFFREYS⁽⁶⁹⁾. The design of a cooler condenser for the case of condensation of two vapours is more

For the condensation of a vapour in the presence of a non-condensable gas, the ollowing example is considered which is based on an the work of KERN⁽²⁸⁾.

Example 9.23

Exchanger consisting of 246 tubes, 19 mm o.d., wall thickness 1.65 mm, 3.65 m long, arranged in four passes soin 25 mm square pitch in a 0.54 m diameter shell and leaves at 322 K. Condensation is effected by cooling water entering and leaving the unit at 300 and 319 K respectively. If the diffusivity of steam-carbon dioxide mixtures is 0.000011 m/s and the group ($\mu_0 D)^{160}$ may be taken to be constant at 0.62, estimate the overall coefficient of heat transfer and the dirt factor for the condenser. Amixture of 0.57 kg/s of steam and 0.20 kg/s of carbon dioxide at 308 kN/m² and its dew point enters a heat

Solution

In the steam entering the condenser, there is
$$\frac{0.57}{18} = 0.032$$
 kmol water
and $\frac{0.20}{44} = 0.0045$ kmol CO₂

Hence the partial pressure of water = $(308 \times 0.032/0.0365) = 270 \text{ kN/m}^2$ and from Table 11A in the Appendix, the dew point = 404 K. Mean molecular weight of the mixture = (0.57 + 0.20)/0.0365 = 21.1 kg/kmol.

At the inlet: vapour pressure of water = 270 kN/m²
inert pressure =
$$(308 - 270) = 38$$
 kN/m²
total = 308 kN/m²

At the outlet: partial pressure of water at 322 K = 11.7 kN/m²
inert pressure =
$$(308 - 11.7) = 296.3 \text{ kN/m}^2$$

steam at the outlet =
$$\frac{0.0045 \times 11.7}{296.3}$$
 = 0.000178 kmol

steam condensed = (0.032 - 0.000178) = 0.03182 kmol.

and:

HEAT TRANSFER

The heat load is now estimated at each interval between the temperatures 404, 401, 397, 380, 339 and

From Table 11A in the Appendix, the partial pressure of steam at 401 $\,\mathrm{K} = 252.2\,\,\mathrm{kN/m^2}$ and hence the partial pressure of $CO_2 = (308 - 252.2) = 55.8 \text{ kN/m}^2$. Steam remaining = $(0.0045 \times 252.2/55.8) = 0.0203 \text{ kmol}$.

 \therefore Steam condensed = (0.032 - 0.0203) = 0.0117 kmol

Heat of condensation = $(0.0117 \times 18)(2180 + 1.93(404 - 401)) = 466 \text{ kW}$

Heat from uncondensed steam = $(0.0203 \times 18 \times 1.93(404 - 401)) = 1.9 \text{ kW}$

Heat from carbon dioxide = $(0.020 \times 0.92(404 - 401)) = 0.5 \text{ kW}$

and the total for the interval = 468.4 kW

Repeating the calculation for the other intervals of temperature gives the following results:

Heat load (kW)	468.4 323.5 343.5 220.1 57.9	1407.3
Interval (K)	404-401 401-397 397-380 380-339 339-322	Total

and the flow of water = $1407.3/(4.187(319 - 300)) = \frac{17.7 \text{ kg/s}}{11.7 \text{ kg/s}}$

With this flow of water and a flow area per pass of 0.0120 m², the mass velocity of water is 1425 kg/m²s, equivalent to a velocity of 1.44 m/s at which $h_i = 6.36 \text{ kW/m}^2 \text{ K}$. Basing this on the outside area $h_{io} = 5.25 \text{ kW/m}^2 \text{ K}.$

Shell-side coefficient for entering gas mixture:

The mean specific heat, $C_p = \frac{(0.20 \times 0.92) + (0.57 \times 1.93)}{0.77} = 1.704 \text{ kJ/kg K.}$

Similarly, the mean thermal conductivity k=0.025 kW/m K and the mean viscosity $\mu=0.015$ mN s/m²

The area for flow through the shell = 0.0411 m^2 and the mass velocity on the shell side

$$= \frac{0.20 + 0.57}{0.0411} = 18.7 \text{ kg/m}^2\text{s}$$

Taking the equivalent diameter as 0.024 m, Re = 29,800

 $h_g = 0.107 \text{ kW/m}^2 \text{ K or } 107 \text{ W/m}^2 \text{ K}.$

$$\left(\frac{\mu}{\rho D}\right)^{0.67} = 0.62, \quad \left(\frac{C_p \mu}{k}\right)^{0.67} = 1.01$$

Now: and:

and:

$$k_G = \frac{h_g(C_p \mu/k)^{0.67}}{C_p P_{sF}(\mu/\rho D)^{0.67}} = \frac{107 \times 1.01}{1704 P_{sF} \times 0.62}$$
$$= \frac{0.102}{P_{sF}}$$

At point 1

Temperature of the gas T = 404 K, partial pressure of steam $P_g = 270 \text{ kN/m}^2$, partial pressure of the from $\Delta_r = 38$ kN/m², water temperature $T_w = 319$ K and $\Delta_T = (404 - 319) = 85$ K. An estimate is now made for the temperature of the condensate film of $T_c = 391$ K. In this case $P_s = 185.4$ kN/m² and $P_s = 185.4$ $(308 - 185.4) = 122.6 \text{ kN/m}^2$.

$$P_{sF} = \frac{122.6 - 38}{\ln(122.6/38)} = 72.2 \text{ kN/m}^2.$$

n, equation 9.181:

$$h_g(T_s - T_c) + k_G \lambda(P_g - P_s) = h_{i0}(T_c - T_{cm})$$

$$0.107(404 - 391) + \left(\frac{0.102}{724}\right) 2172(270 - 185.4) = 5.25(391 - 319)$$

i.e. there is no balance 259 = 378

$$\prod_{i=1}^{4} y_i T_c = 378 \text{ K}, P_s = 118.5 \text{ kN/m}^2, P_g = (308 - 118.5) = 189.5 \text{ kN/m}^2$$

$$P_s F = \frac{189.5 - 38}{\ln(189.5/38)} = 94.2 \text{ kN/m}^2$$
and:

$$0.107(404 - 378) + \left(\frac{0.102}{94.2}\right) 2172(270 - 118.5) = 5.25(378 - 319)$$

$$U\Delta T = 309 \text{ kW/m}^2$$
 and $U = \frac{309}{(404 - 319)}$

310 = 308 which agrees well

Repeating this procedure at the various temperature points selected, the heat-exchanger area may then be obtained as the area under a plot of Σq vs. $1/U\Delta T$, or as $A = \Sigma q/U\Delta T$ according to the following tabulation: $= 3.64 \text{ kW/m}^2 \text{ K}.$

Point	T _s (K)	T _c (K)	UAT (kW/m²)	$(V\Delta T)_{ow}$ (kW/m^2)	(kW)	$A = Q/(U\Delta T)_{ow}$ (m ²)	Δ <i>T</i> (K)	Δ <i>T</i> σ _w (K)	$Q/\Delta T_{ow}$ (kW/K)
1	\$	378	309	1		1	84.4	ı	ı
75	6	326	228	268.5	468.4	1.75	88.1	86.3	5.42
33	397	336	145	186.5	323.5	1.74	88.6	88.4	3.66
4	380	312	40.6	88.1*	343.5	3.89	76.7	82.7	4.15
S.	339	302	5.4	17.5*	220.1	12.58	38.1	55.2*	4.00
9	322	300	2.1	3.5*	51.9	14.83	22.2	29.6*	1.75
				Total:	1407.3	34.8			18.98

*based on LMTD.

 $\frac{1}{2}$ If no condensation takes place, the logarithmic mean temperature difference is 46.6 K. In practice the value $\frac{1}{2}(1407.3/18.98) = 74.2$ K.

Assuming no scale resistance, the overall coefficient is $\frac{1407.3}{34.8 \times 74.2} = 0.545 \text{ kW/m}^2 \text{ K}$.

he available surface area on the outside of the tubes $= 0.060 \text{ m}^2/\text{m}$

or $(246 \times 3.65 \times 0.060) = 53.9 \text{ m}^2$

The actual coefficient is therefore $\frac{1407.5}{53.9 \times 74.2} = 0.352 \text{ kW/m}^2 \text{ K}$ 1407.3

As shown in Figure 9.50, the clean coefficient varies from $3.64 \text{ kW/m}^2\text{ K}$ at the inlet to $0.092 \text{ kW/m}^2\text{ K}$ at $\frac{1}{1000} = \frac{1}{1000} = \frac{1}{100$



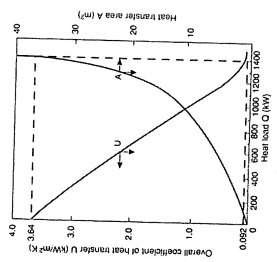


Figure 9.50. Results for Example 9.23

Condensation of mixed vapours is considered further in Volume 6, Chapter 12, where it is suggested that the local heat transfer coefficient may be expressed in terms of the local gas-film and condensate-film coefficients. For partial condensation where:

- non-condensables < 0.5 per cent; their effect can be ignored,
- (ii) non-condensables > 70 per cent; the heat transfer can be taken as being by forced convection alone, and
 - (iii) non-condensables 0.5-70 per cent; both mechanisms are effective.

9.7. BOILING LIQUIDS

9.7.1 Conditions for boiling

In processing units, liquids are boiled either on submerged surfaces or on the inside The boiling of liquids under either of these conditions normally leads to the formation of vertical tubes. Mechanical agitation may be applied in the first case and, in the second, the liquid may be driven through the tubes by means of an external pump of vapour first in the form of bubbles and later as a distinct vapour phase above. liquid interface. The conditions for boiling on the submerged surface are discussed here and the problems arising with boiling inside tubes are considered in Volume 2, Much of the fundamental work on the ideas of boiling has been presented by WESTWATTER (19) and $\mathsf{JAKOB}^{(71)}$, and subsequently by ROHSENOW and CLARK $^{(72)}$ and ROHSENOW $^{(33)}$ and $b^{s_0}_2$

HEAT TRANSFER

ENSTER (74). The boiling of solutions in which a solid phase is separated after evaporation has proceeded to a sufficient extent is considered in Volume 2.

the surface tension between the liquid and vapour, the vapour pressure on the inside of a oncave surface will be less than that at a plane surface. As a result, the vapour pressure For a bubble to be formed in a liquid, such as steam in water, for example, it is necessary for a surface of separation to be produced. Kelvin has shown that, as a result of $p_i^{\rm s}$ inside the bubble is less than the saturation vapour pressure P_i at a plane surface. The relation between P, and P, is:

$$P_r = P_s - \left(\frac{2\sigma}{r}\right) \tag{9.184}$$

where r is the radius of curvature of the bubble, and σ is the surface tension.

that very small bubbles are difficult to form without excessive superheat. The formation of bubbles is made much easier by the fact that they will form on curved surfaces or on aperheat increasing with decrease in the radius of the bubble. On this basis it follows Hence the liquid must be superheated near the surface of the bubble, the extent of the pregularities on the heating surface, so that only a small degree of superheat is normally required.

most promising. In many cavities the angle θ will be greater than 90° and the effective Nucleation at much lower values of superheat is believed to arise from the presence of existing nuclei such as non-condensing gas bubbles, or from the effect of the shape of the cavities in the surface. Of these, the current discussion on the influence of cavities is the confact angle, which includes the contact angle of the cavity β , will be considerably greater $=\theta + (180 - \beta)/21$, so that a much-reduced superheat is required to give nucleation. Thus the size of the mouth of the cavity and the shape of the cavity plays a significant

part in nucleation⁽⁷⁵⁾.

It follows that for boiling to occur a small difference in temperature must exist between flux of about 25.2 kW/nr2, a temperature difference of only 6 deg K was required with a the liquid and the vapour. JAKOB and FRITZ. (76) have measured the temperature distribution for water boiling above an electrically heated hot plate. The temperature dropped very steeply from about 383 K on the actual surface of the plate to 374 K about 0.1 mm from it. Beyond this point the temperature was reasonably constant until the water surface was ractical. The mean superheat of the water above the temperature in the vapour space was about 0.5 deg K and this changed very little with the rate of evaporation. At higher pressures this superheating became smaller becoming 0.2 deg K at 5 MN/m² and 0.05 deg K at 101 MN/m2. The temperature drop from the heating surface depends, however, very much rough surface as against 10.6 deg K with a smooth surface. The heat transfer coefficient on the boiling side is therefore dependent on the nature of the surface and on the difference in temperature available. For water boiling on copper plates JAKOB and FRITZ[76] give the following coefficients for a constant temperature difference of 5.6 deg K, with different on the rate of heat transfer and on the nature of the surface. Thus in order to maintain a heat

- (1) Surface after 8 h (28.8 ks) use and 48 h (172.8 ks)
 - immersion in water
 - (2) Freshly sandblasted
- (3) Sandblasted surface after long use
- $h = 8000 \text{ W/m}^2 \text{ K}$
- $h = 3900 \text{ W/m}^2 \text{ K}$ $h = 2600 \text{ W/m}^2 \text{ K}$

HEAT TRANSFER

(4) Chromium plated

$$h = 2000 \text{ W/m}^2 \text{ K}$$

The initial surface, with freshly cut grooves, gave much higher figures than case (1). The nature of the surface will have a marked effect on the physical form of the bubble and the area actually in contact with the surface, as shown in Figure 9.51.

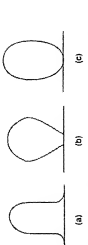


Figure 9.51. Shapes of bubbles (a) screen surface—thin oil layer (b) chromium plated and polished surface—(c) screen surface—clean

The three cases are:

- (a) Non-wettable surface, where the vapour bubbles spread out thus reducing the area available for heat transfer from the hot surface to the liquid.
 - (b) Partially wettable surface, which is the commonest form, where the bubbles rise from a larger number of sites and the rate of transfer is increased.
- (c) Entirely wented surface, such as that formed by a screen. This gives the minimum area of contact between vapour and surface and the bubbles leave the surface when still very small. It therefore follows that if the liquid has detergent properties this may give rise to much higher rates of heat transfer.

9.7.2. Types of boiling

Interface evaporation

In boiling liquids on a submerged surface it is found that the heat transfer coefficient depends very much on the temperature difference between the hot surface and the boiling liquid. The general relation between the temperature difference and heat transfer coefficient was first presented by NUKIYAMA⁽⁷⁷⁾ who boiled water on an electrically heated wire. The results obtained have been confirmed and extended by others, and Figure 9.53 shows the data of FARBER and SCORAH⁽⁷⁸⁾. The relationship here is complex and is best considered in stages.

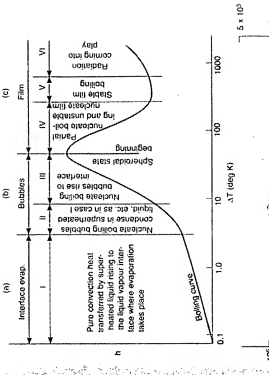
In interface evaporation, the bubbles of vapour formed on the heated surface move to the vapour-liquid interface by natural convection and exert very little agitation on the liquid. The results are given by:

$$Nu = 0.61(Gr Pr)^{1/4} (9.185)$$

which may be compared with the expression for natural convection:

$$Nu = C'(Gr Pr)^n (equation 9.101)$$

where n = 0.25 for streamline conditions and n = 0.33 for turbulent conditions,



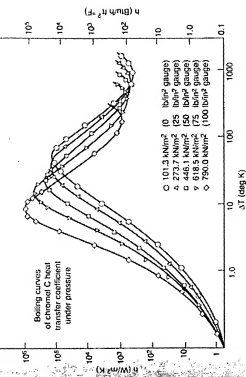


Figure 9.52. Heat transfer results of FARBER and SCURAH⁽⁷⁸⁾

Nucleate boiling

At higher values of ΔT the bubbles form more rapidly and form more centres of furtheration. Under these conditions the bubbles exert an appreciable agitation on the liquid and the heat transfer coefficient rises rapidly. This is the most important region for boiling the infinite equipment.

HEAT TRANSFER

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Film boiling

With a sufficiently high value of ΔT, the bubbles are formed so rapidly that they cannon means that the liquid is prevented from flowing on to the surface by the bubbles of vapour and the coefficient falls. The maximum coefficient occurs during nucleate boiling although this is an unstable region for operation. In passing from the nucleate boiling region to itself in a decrease in the heat flux, the second is the prelude to stable film boiling. The first change in the process. The first manifests intermediate region is generally known as the transition region. It may be noted that the first change in the process is an important hydrodynamic phenomenon which is common systems, for example.

With very high values of ΔT , the heat transfer coefficient rises again because of heat transfer by radiation. These very high values are rarely achieved in practice and usually the aim is to operate the plant at a temperature difference a little below the value giving the maximum heat transfer coefficient.

9.7.3. Heat transfer coefficients and heat flux

The values of the heat transfer coefficients for low values of temperature difference are given by equation 9.185. Figure 9.53 shows the values of h and for q for water boiling on a submerged surface. Whilst the actual values vary somewhat between investigations they all give a maximum for a temperature difference of about 22 deg K. The maximum value of h is about 50 kW/m² K and the maximum flux is about 1100 kW/m².

Similar results have been obtained by BONILLA and PERRY⁽⁷⁹⁾, INSINGER and BLISS⁽⁸⁰⁾ and others for a number of organic liquids such as benzene, alcohols, acctone, and Carbon tetrachloride. The data in Table 9.9 for liquids boiling at atmospheric pressure show that the maximum heat flux is much smaller with organic liquids than with water and the temperature difference at this condition is rather higher. In practice the critical value of AT may be exceeded. SAUER et al. (81) found that the overall transfer coefficient U for steam pressure was reduced to 115 kN/m² was only 14 per cent of that when the

Table 9.9. Maximum heat flux for various liquids boiling at atmospheric pressure

		The state of the s	
Lìquid	Surface	Critical AT (deg K)	Maximum flux (kW/m²)
Water 50 mol/s ethanol-water Ethanol n-Butanol iso-Butanol Acetione iso-Proganol Carbon tetrachloride Benzene	Chromium Chromium Chromium Chromium Nickel Chromium Chromium Copper Copper	888448811	910 595 455 455 370 340 180
The state of the s	Constitution of the last of th		Charles Co.

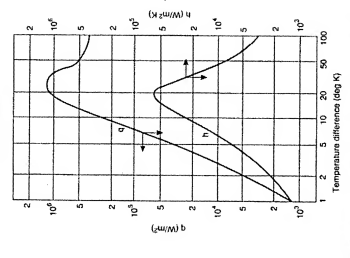


Figure 9.53. Effect of temperature difference on heat flux and heat transfer coefficient to water boiling at 373 K on a submerged surface

In considering the problem of nucleate boiling, the nature of the surface, the pressure, and the temperature difference must be taken into account as well as the actual physical properties of the liquid.

Apart from the question of scale, the nature of the clean surface has a pronounced influence on the rate of boiling. Thus BONILLA and PERRY⁽⁷⁹⁾ boiled ethanol at atmospheric pressure and a temperature difference of 23 deg K, and found that the heat flux atamospheric pressure was 850 kW/m² for polished copper, 450 for gold plate, and 370 for fresh chromium plate, and only 140 for old chromium plate. This wide fluctuation means that care must be taken in anticipating the heat flux, since the high values that may be obtained initially may not persist in practice because of tarnishing of the surface.

Effect of temperature difference

CRYDER and FINALBORGO⁽⁸²⁾ boiled a number of liquids on a horizontal brass surface, both at atmospheric and at reduced pressure. Some of their results are shown in Figure 9.54, where the coefficient for the boiling liquid h is plotted against the temperature difference between the hot surface and the liquid. The points for the various liquids in Figure 9.54

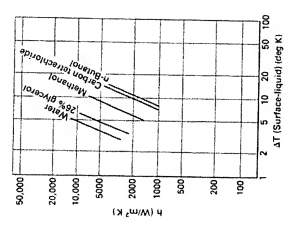


Figure 9.54. Effect of temperature difference on the heat transfer coefficient for boiling liquids (Chypek and First, RORGO^(8.2))

lie on nearly parallel straight lines, which may be represented by:

$$h = \text{constant} \times \Delta T^{2.5} \tag{9.186}$$

This value for the index of ΔT has been found by other workers, although JAKOB and LINKE⁽⁸³⁾ found values as high as 4 for some of their work. It is important to note that this value of 2.5 is true only for temperature differences up to 19 deg K.

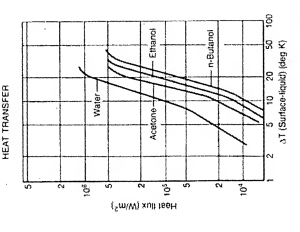
In some ways it is more convenient to show the results in the form of heat flux versus temperature difference, as shown in Figure 9.55, where some results from a number of workers are given.

Effect of pressure

CRYDER and FINALBORGO⁽⁸²⁾ found that h decreased uniformly as the pressure and hence the boiling point was reduced, according to the relation $h = \text{constant} \times B^{T'}$, where T' is numerically equal to the temperature in K and B is a constant. Combining this with equation 9.186, their results for h were expressed in the empirical form:

or, using SI units:
$$\log\left(\frac{h}{5.67}\right) = a' + 2.5 \log \Delta T + b'(T'' - 273)$$
 (9.187)

where (T'' - 273) is in °C.



gure 9.55. Effect of temperature difference on heat flux to boiling liquids (Bonn), and Perry (20)

If a and b' are given the following values, h is expressed in Wtm^2K :

	ϫ	æ		a,	P,
Water	-0.96	0.025	Kerosene	-4.13	0.022
Methanol	*****	0.027	10% Na ₂ SO ₃	-1.47	0.029
CCL	-1.55	0.022	24% NaCl	-2.43	0.031

The values of a' will apply only to a particular apparatus although a value of b' of 0.025 is of more general application. If h_n is the coefficient at some standard boiling point T_n and b' at some other temperature T, equation 9.187 may be rearranged to give:

$$\log \frac{h}{h_n} = 0.025(T^n - T_n^n)$$
 (9.188)

for a given material and temperature difference.

As the pressure is raised above atmospheric pressure, the film coefficient increases for a constant temperature difference. CICHELLI and BONILLA⁽⁸⁴⁾ have examined this problem for pressures up to the critical value for the vapour, and have shown that ΔT for maximum rate of boiling decreases with the pressure. They obtained a single curve, shown in Figure 9.56, by plotting q_{max}/P_c against P_R , where P_c is the critical pressure and P_R the reduced pressure = P/P_c . This curve represents the data for water, ethanol. because, propane, n-heptane, and several mixtures with water. For water the results cover only a small range of P/P_c , because of the high value of P_c . For the organic liquids



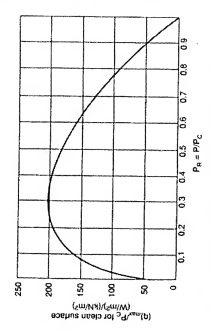


Figure 9.56. Effect of pressure on the maximum heat flux in nucleate boiling

P of about one-third of the critical pressure P_c . As shown in Table 9.10, the range of physical properties of the organic liquids is not wide and further data are required to investigated, it was shown that the maximum value of heat flux q occurs at a pressure substantiate the previous relation.

Table 9, 10, Heat transfer

	Boiling	ΔŢ		ħ
Liquid	(deg K)	(deg K)	(W/m2 K)	(Bru/h fr ² °F)
Water	372	4.7	0XX6	1600
	372	2.9	2700	200
	326	90°	4700	850
	326	6.1	1300	250
Methanol	337	8.9	4800	850
	337	5.6	1500	250
	306	14.4	3000	200
	308	9.3	900	150
Carbon tetrachlorida	349	12.6	3500	99
	9X9	7.2	90=	200
	315	70.	2000	9
	315	8; =	700	25

9.7.4. Analysis based on bubble characteristics

It is a matter speculation as to why such high values of heat flux are obtained with the which was added to the liquid by their movement. It has now been shown, by determining part of the heat that is actually transferred. The current views are that the high flux arises the numbers of bubbles, that this mechanism would result in the transfer of only a moderate boiling process. It was once thought that the bubbles themselves were carriers of latent heal

HEAT TRANSFER

from the agitation produced by the bubbles, and two rather different explanations have been put forward. ROHSENOW and CLARK $^{(72)}$ and ROHSENOW $^{(33)}$ base their argument on size of the bubble an expression may be derived for the heat transfer coefficient in the form of a Nusselt type equation, relating the Nusselt group to the Reynolds and Prandtl goups. FORSTER and ZUBER^(85,86), however, argue that the important velocity is that of he growing bubble, and this is the term used to express the velocity. In either case the bubble movement is vital in obtaining a high flux. The liquid adjacent to the surface is aginated and exerts a mixing action by pushing hot liquid from the surface to the bulk of decondition of the bubble on leaving the hot surface. By calculating the velocity and the stream.

Considering in more detail the argument proposed by ROHSENHOW and CLARK (72) and ROHSENHOW (73), the size of a bubble at the instant of breakaway from the surface has determined by FRITZ⁽⁸⁷⁾ who has shown that d_b is given by:

$$d_b = C_1 \phi \left(\frac{2\sigma}{g(\rho_1 - \rho_v)} \right)^{1/2} \tag{9.189}$$

where σ is the surface tension, ρ_l and ρ_v the density of the liquid and vapour, ϕ is the contact angle, and C1 is a constant depending on conditions.

The flowrate of vapour per unit area as bubbles up is given by:

$$u_b = \frac{f_H \pi d_b^3}{6} \tag{9.190}$$

where f is the frequency of bubble formation at each bubble site and n is the number of sites of nucleation per unit area.

The heat transferred by the bubbles q, is to a good approximation given by:

$$q_b = \frac{1}{\delta}\pi d_b^3 f n \rho_v \lambda \tag{9.191}$$

where A is the latent heat of vaporisation.

It has been shown that for heat flux rates up to 3.2 kW/m² the product fd_b is constant indition the total heat flow per unit area q is proportional to n. From equation 9.191 it is seen that q_b is proportional to n at a given pressure, so that $q \propto q_b$.

$$q = C_2 \frac{\pi}{6} d_b^3 f n \rho_b \lambda \tag{9.192}$$

Hence:

Substituting from equations 9.190 and 9.192, the mass flow per unit area:

$$\rho_{\nu}u_{b} = fn\frac{\pi}{6}d_{b}^{3}\rho_{\nu} = \frac{q}{C_{2}\lambda}$$
 (9.193)

A Reynolds number for the bubble flow which represents the term for agitation may be defined as:

$$Re_b = \frac{d_b \rho_u u_b}{\mu_1}$$

$$= C_1 \phi \left(\frac{2\sigma}{g(\rho_1 - \rho_v)} \right)^{1/2} \left(\frac{q}{C_2 \lambda} \right)_1$$

492

$$=C_3\phi\frac{q}{\lambda\mu_l}\left(\frac{\sigma}{g(\rho_l-\rho_v)}\right)^{1/2}$$

(9.194)

The Nusselt group for bubble flow, $Nu_b = h_b C_1 \frac{\phi}{k_l} \left(\frac{2\sigma}{g(\rho_l - \rho_v)} \right)^{1/2}$

$$=C_4h_b\frac{\phi}{k_l}\left(\frac{\sigma}{8(\rho_l-\rho_v)}\right)^{1/2} \tag{9.195}$$

and hence a final correlation is obtained of the form:

$$Nu_b = \text{constant } Re_b^n Pr^m$$

(9.196)

$$Nu_b = \text{constant} \left[\frac{C_3 \phi q}{\mu l \lambda} \left(\frac{\sigma}{g(\rho_l - \rho_v)} \right)^{1/2} \right]^n \left(\frac{C_I \mu_I}{k_I} \right)^m \tag{9.197}$$

Ċ.

where n and m have been found experimentally to be 0.67 and -0.7 respectively and the constant, which depends on the metal surface, ranges from 67–100 for polished chromium, 77 for platinum wire and 166 for brass. (73)

A comprehensive study of nucleate boiling of a wide range of liquids on thick plates of copper, aluminium, brass and stainless steel has been carried out by PiORO^(SS) who has evaluated the constants in equation 9,197 for different combinations of liquid and surface.

FORSTER and ZUBER^(83,86) who employed a similar basic approach, although the radial rate of growth dr/dr was used for the bubble velocity in the Reynolds group, showed that:

$$\frac{dr}{dt} = \frac{\Delta T C_I \rho_i}{2\lambda \rho_c} \left(\frac{\pi D_{HI}}{t} \right)^{1/2} \tag{9.198}$$

where D_{Hl} is the thermal diffusivity $(k_I/C_I\rho_I)$ of the liquid. Using this method, a final correlation in the form of equation 9.196 has been presented.

Although these two forms of analysis give rise to somewhat similar expressions, the basic terms are evaluated in quite different ways and the final expressions show many differences. Some data fit the Rohsenow equation reasonably well⁽⁸⁸⁾, and other data fit Forster's equation.

These expressions all indicate the importance of the bubbles on the rate of transfer, although as yet they have not been used for design purposes. Insinger and BLISs⁽⁸⁰⁾ made the first approach by dimensional analysis and MCNELLY⁽⁸⁹⁾ has subsequently obtained a more satisfactory result. The influence of ΔT is taken into account by using the flux q_s and the last term allows for the change in volume when the liquid vaporises. The following expression was obtained in which the numerical values of the indices were deduced from existing data:

$$\frac{hd}{k_l} = 0.225 \left(\frac{C_l \mu_l}{k_l} \right)^{0.09} \left(\frac{qd}{\lambda \mu} \right)^{0.09} \left(\frac{Pd}{\sigma} \right)^{0.31} \left(\frac{\rho_l}{\rho_o} - 1 \right)^{0.33}$$
 (9.199)

9.7.5. Sub-cooled boiling

If bubbles are formed in a liquid which is much below its boiling point, then the bubbles will collapse in the bulk of the liquid. Thus if a liquid flows over a very hot surface; then

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the bubbles formed are carried away from the surface by the liquid and sub-cooled boiling ocurs. Under these conditions a very large number of small bubbles are formed and a very high heat flux is obtained. Some results for these conditions are given in Figure 9.57.

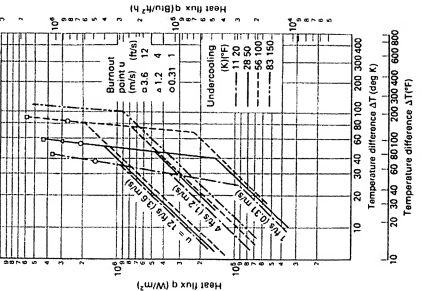


Figure 9.57. Heat flux in sub-cooled boiling

If a liquid flows through a tube heated on the outside then the heat flux q will increase with ΔT as shown in Figure 9.57. Beyond a certain value of ΔT the increase in q is very appear in the velocity through the tube is increased, then a similar plot is obtained with a ligher value of q at low values of ΔT and then the points follow the first line. Over the fix section, forced convection boiling exists where an increase in Reynolds number does not bring about a very great increase in q because the bubbles are themselves producing adiation in the boundary layer near the wall. Over the steep section, sub-cooled boiling exists where the velocity is not important provided it is sufficient to remove the bubbles

HEAT TRANSFER

rapidly from the surface. In the same way, mechanical agitation of a liquid boiling on a submerged surface will not markedly increase the heat flux.

9.7.6. Design considerations

In the design of vaporisers and reboilers, two types of boiling are important—nucleate boiling in a pool of liquid as in a kettle-type reboiler or a jacketed vessel, and convective boiling which occurs where the vaporising liquid flows over a heated surface and heat transfer is by both forced convection and nucleate boiling as, for example, in forced circulation or thermosyphon reboilers. The discussion here is a summary of that given in Volume 6 where a worked example is given.

In the absence of experimental data, the correlation given by FORSTER and ZUBER(⁸⁶⁾ may be used to estimate *pool boiling* coefficients, although the following reduced pressure correlation given by MOSTINSKI⁽⁹⁰⁾ is much simpler to use and gives reliable results for h (in W/m² K):

$$h = 0.104 P_c^{0.69} q^{0.7} \left[1.8 \left(\frac{P}{P_c} \right)^{0.17} + 4 \left(\frac{P}{P_c} \right)^{1.2} + 10 \left(\frac{P}{P_c} \right)^{10} \right]$$
 (9.200)

In this equation, P_c and P are the critical and operating pressures (bar), respectively, and q is the heat flux (W/m^2). Both equations are for single component fluids, although they may also be used for close-boiling mixtures and for wider boiling ranges with a factor of safety. In reboiler and vaporiser design, it is important that the heat flux is well below the critical value. A correlation is given for the heat transfer coefficient for the case where film-boiling takes place on tubes submerged in the liquid.

Convective boiling, which occurs when the boiling liquid flows through a tube or over a tube bundle, depends on the state of the fluid at any point. The effective heat transfer coefficient can be considered to be made up of the convective and nucleate boiling components. The convective boiling coefficient is estimated using an equation for single-phase forced-convection heat transfer (equation 9.64, for example) modified by a factor to allow for the effects of two-phase flow. Similarly, the nucleate boiling coefficient is obtained from the Forster and Zuber or Mostinski correlation, modified by a factor dependent on the liquid Reynolds number and on the effects of two-phase flow. The estimation of convective boiling coefficients is illustrated by means of an example in Volume 6.

One of the most important areas of application of heat transfer to boiling liquids is in the use of evaporators to effect an increase in the concentration of a solution. This topic is considered in Volume 2.

For vaporising the liquid at the bottom of a distillation column a reboiler is used, as shown in Figure 9.58. The liquid from the still enters the boiler at the base, and, after flowing over the tubes, passes out over a weir. The vapour formed, together with any entrained liquid, passes from the top of the unit to the column. The liquid flow may be either by gravity or by forced circulation. In such equipment, provision is made for expansion of the tubes either by having a floating head as shown, or by arranging the tubes in the form of a hairpin bend (Figure 9.59). A vertical reboiler may also be used with steam condensing on the outside of the tube bundle. With all systems it is undesirable

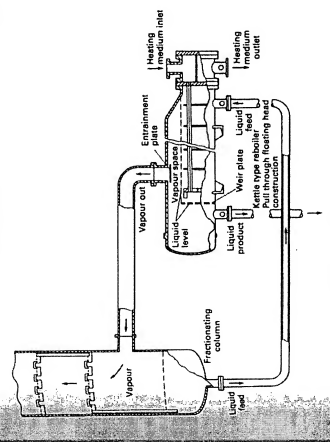


Figure 9.58. Reboiler installed on a distillation column

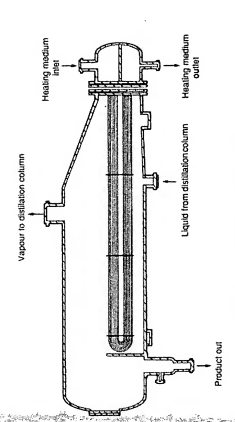


Figure 9.59. Kettle rehoiler with hairpin tubes

to vaporise more than a small percentage of the feed since a good liquid flow over the tubes is necessary to avoid scale formation.

In the design of farced canvection reboilers, the normal practice is to calculate the head an assumed circulation rate through the exchanger. Kettle reboilers, such as that shown 60 kW/m2 for organics and 90 kW/m2 for dilute aqueous solutions. In thermosyphonic balanced by the hydrostatic head and the design involves an iterative procedure based on in Figure 9.59, are essentially pool boiling devices and their design, based on nucleage boiling data, uses the Zuber equation for single tubes, modified by a tube-density facing and this gives safe values. KERN(28) recommends that the heat flux should not exceed rebuilers, the fluid circulates at a rate at which the pressure losses in the system are just transfer coefficient on the assumption that heat is transferred by forced convection only This general approach is developed further in Volume 6.

9.8. HEAT TRANSFER IN REACTION VESSELS

9.8.1. Helical cooling coils

A simple jacketed pan or kettle is very commonly used in the processing industries a reaction vessel. In many cases, such as in nitration or sulphonation reactions, heat has bring it to completion. The addition or removal of heat is conveniently arranged in passing steam or water through a jacket fitted to the outside of the vessel or through helical coil fitted inside the vessel. In either case some form of agitator is used to obrain even distribution in the vessel. This may be of the anchor type for very thick pastes or a to be removed or added to the mixture in order either to control the rate of reaction; or propeller or turbine if the contents are not too viscous.

In such a vessel, the thermal resistances to heat transfer arise from the water film on the inside of the coil, the wall of the tube, the film on the outside of the coil, and any scale that may be present on either surface. The overall transfer coefficient may be expressed by

$$\frac{1}{UA} = \frac{1}{h_i A_i} + \frac{x_w}{k_w A_w} + \frac{1}{h_o A_o} + \frac{R_o}{A_o} + \frac{R_i}{A_i}$$
 (9.201)

where R_o and R_i are the scale resistances and the other terms have the usual definitions

Inside film coefficient

The value of h_i may be obtained from a form of equation 9.64:

$$\frac{h_i d}{k} = 0.023 \left(\frac{du\rho}{\mu}\right)^{0.3} \left(\frac{C_{\mu}\mu}{k}\right)^{0.33} \tag{9.202}$$

if water is used in the coil, and the Sieder and Tate equation (equation 9.66) if a viscous brine is used for cooling.

These equations have been obtained for straight tubes; with a coil somewhat greater transfer is obtained for the same physical conditions. JESCHKE(91) cooled air in a 31 mm steel tube wound in the form of a helix and expressed his results in the form:

$$h_i(\text{coil}) = h_i(\text{straight pipe}) \left(1 + 3.5 \frac{d}{d_c}\right)$$
 (9.203)

HEAT TRANSFER

Let d is the inside diameter of the tube and d_c the diameter of the helix. PRATE⁽⁹²⁾ has Examined this problem in greater detail for liquids and has given almost the same result. Combining equations 9.202 and 9.203, the inside film coefficient h_i for the coil may be calculated.

Toutside film coefficient

The value of h_o is determined by the physical properties of the liquor and by the degree of agitation achieved. This latter quantity is difficult to express in a quantitative manner and the group $L^2N\rho/\mu$ has been used both for this problem and for the allied one of power used in agitation, as discussed in Chapter 7. In this group L is the length of the gaddle and N the revolutions per unit time. CHILTON, DREW and JEBENS⁽⁹³⁾, working with Small tank only 0.3 m in diameter d_{ν} , expressed their results by:

$$\frac{h_o d_v}{k} \left(\frac{\mu_x}{\mu}\right)^{0.14} = 0.87 \left(\frac{C_p \mu}{k}\right)^{1/3} \left(\frac{L^2 N \rho}{\mu}\right)^{0.62} \tag{9.204}$$

where the factor $(\mu_s/\mu)^{0.14}$ allows for the difference between the viscosity adjacent to The coil (μ_s) and that in the bulk of the liquor. A wide range of physical properties was satisfieved by using water, two oils, and glycerol.

PRATT (92) used both circular and square tanks up to 0.6 m in size and a series of different arrangements of a simple paddie as shown in Figure 9.60. The effect of altering the arrangement of the coil was investigated and the tube diameter d_o , the gap between the turns d_g , the diameter of the helix d_c , the height of the coil d_ρ , and the width of the surrect W were all varied. The final equations tanks were:

For cylindrical tanks

(9.205)

$$\frac{h_o d_v}{k} = 34 \left(\frac{L^2 N \rho}{\mu} \right)^{0.5} \left(\frac{d_g}{k} \right)^{0.3} \left(\frac{W}{d_c} \right)^{0.25} \left(\frac{L^2 d_v}{d_d^3} \right)^{0.11}$$

Figure 9.60. Arrangement of coil in Prair's work (79)

For square tanks:

$$\frac{h_0 I_v}{k} = 39 \left(\frac{L^2 N \rho}{\mu} \right)^{0.5} \left(\frac{C_{\rho \mu}}{k} \right)^{0.3} \left(\frac{d_g}{d_{\rho}} \right)^{0.8} \left(\frac{W}{d_c} \right)^{0.25} \left(\frac{L^2 I_v}{d_s^3} \right)^{0.1}$$
(9.2)

where I_v is the length of the side of the vessel.

conditions. CUMMINGS and WEST⁽⁹⁴⁾ have tested these results with a much larger tank of 0.45 m³ capacity and have given an expression similar to equation 9.204 but with a many cases a second impeller was mounted above the first, giving an agitation which is probably more intense than that attained by the other workers. A constant of 0.9 seems a These give almost the same results as the earlier equations over a wide range of constant of 1.01 instead of 0.87. A retreating blade turbine impeller was used, and in reasonable average from existing work.

Example 9.24

Toluene is continuously nitrated to monomitrotoluene in a cast-iron vessel, 1 m diameter, fitted with a propellicr agitator 0.3 m diameter rotating at 2.5 Hz. The temperature is maintained at 310 K by circulating 0.5 kgs cooling water through a stainless steel coil 25 mm o.d. and 22 mm i.d. wound in the form of a helix, 0.80 m in diameter. The conditions are such that the reacting material may be considered to have the same physical properties as 75 per cem sulphune acid. If the mean water temperature is 290 K, what is the overall coefficient of heat transfer?

The overall coefficient Uo based on the outside area of the coil is given by equation 9.201:

$$\frac{1}{U_0} = \frac{1}{h_v} + \frac{s_w d_o}{k_w d_w} + \frac{d_o}{h_i d} + R_o + \frac{R_i d_o}{d}$$

where d., is the mean diameter of the pipe, From equations 9.202 and 9.203, the inside film coefficient for the water is given by:

$$h_i = \frac{k}{d} \left(1 + 3.5 \frac{d}{d_e} \right) 0.023 \left(\frac{dup}{\mu} \right)^{0.8} \left(\frac{C_p \mu}{k} \right)^{0.4}$$

$$\rho u = \frac{0.5}{(\pi/4) \times 0.022^2} = 1315 \text{ kg/m}^2 \text{s}$$

d = 0.022 m, $d_{\rm r} = 0.80$ m, k = 0.59 W/m K, $\mu = 1.08$ mN s/m² or 1.08×10^{-3} N s/m², and $C_{\rm p} = 4.18 \times 10^3$ J/kg K

Thus:
$$h_i = \frac{0.59}{0.022} \left(1 + 3.5 \times \frac{0.022}{0.80} \right) 0.023 \left(\frac{0.022 \times 1315}{1.08 \times 10^{-3}} \right)^{0.8} \left(\frac{4.18 \times 10^3 \times 10^{-3}}{0.59} \right)^{0.4}$$

 $= 0.680(26.780)^{0.8}(7.65)^{0.4} = 5490 \text{ W/m}^2 \text{ K}$

The external film coefficient is given by equation 9.204:

$$\frac{h_0 d_n}{k} \left(\frac{\mu_4}{\mu} \right)^{0.14} = 0.87 \left(\frac{C_P \mu}{k} \right)^{0.33} \left(\frac{L^2 N \rho}{\mu} \right)^{0.62}$$

For 75 per cent sulphuric acid;

HEAT TRANSFER

 $L_{\rm eff} = 0.40$ Whn K, $\mu_s = 8.6 \times 10^{-3}$ Ns/m² at 300 K, $\mu = 6.5 \times 10^{-3}$ Ns/m² at 310 K, $C_p = 1.88 \times 10^3$ JAS K. and p = 1666 kg/m

$$\frac{h_0 \times 1.0}{0.40} \left(\frac{8.6}{6.5}\right)^{0.14} = 0.87 \left(\frac{1.88 \times 10^3 \times 6.5 \times 10^{-3}}{0.40}\right)^{0.35} \left(\frac{0.3^2 \times 2.5 \times 1666}{6.5 \times 10^{-3}}\right)^{0.62}$$

 $2.5h_o \times 1.04 = 0.87 \times 3.09 \times 900$ $h_o = 930 \text{ W/m}^2 \text{ K}$

Taking
$$k_0^2=15.9$$
 W/m K and R_0 and R_1 as 0.0004 and 0.0002 m² K/W, respectively:

$$\frac{1}{U_o} = \frac{1}{930} + \frac{0.0015 \times 0.025}{15.9 \times 0.0235} + \frac{0.025}{5490 \times 0.022} + 0.0004 + \frac{0.0002 \times 0.025}{0.022} = 0.00107 + 0.00010 + 0.00021 + 0.00040 + 0.00023 = 0.00201$$

 $U_o = 498 \text{ W/m}^2 \text{ K}$

fights calculation a mean area of surface might have been used with sufficient accuracy. It is important to not the importance of the scale terms which together form a major part of the thermal resistance,

9.8.2. Jacketed vessels

by condensing steam in a jacket or passing water through it - an arrangement which is often used for organic reactions where the mixture is too viscous for the use of coils and a high-speed agitator. CHILTON et al. (93) and CUMMINOS and WEST (94) have measured the hear is supplied to the jacket and simultaneously removed by passing water through the and the surface of the coil by means of thermocouples and thus obtained the film heat transfer coefficients for this case by using an arrangement as shown in Figure 9.61, where transfer coefficients directly. Cummings and West used an indirect method to give the In many cases, heating or cooling of a reaction mixture is most satisfactorily achieved coil Chilton measured the temperatures of the inside of the vessel wall, the bulk liquid, film coefficient from measurements of the overall coefficients.

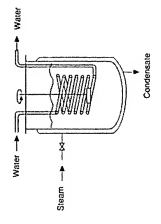


Figure 9.61. Reaction vessel with jacket and coil

CHILTON et al. (93) expressed their results by:

$$\frac{h_b d_v}{k} \left(\frac{\mu_s}{\mu}\right)^{0.14} = 0.36 \left(\frac{L^2 N \rho}{\mu}\right)^{0.67} \left(\frac{C_\rho \mu}{k}\right)^{0.33} \tag{9.207}$$

where h_b is the film coefficient for the liquor adjacent to the wall of the vessel. CUMMINGS and WEST^(9,1) used the same equation although the coefficient was 0.40. Considering final Chilton's vessel was only 0.3 m in diameter and fitted with a single paddle of 150 mm length, and that Cummings and West used a 0.45 m³ vessel with two turbine impellers agreement between their results is remarkably good. The group $(\mu_s/\mu)^{0.14}$ is again used to allow for the difference in the viscosities at the surface and in the bulk of the fluid.

BROWN et al. (95) have given data on the performance of 1.5 m diameter sulphonator, and nitrators of 3.4 m³ capacity as used in the dyestuffs industry. The sulphonators were of cast iron and had a wall thickness of 25.4 mm; the annular space in the jacket being also 25.4 mm. The agitator of the sulphonator was of the anchor type with a 127 mm clearance at the walls and was driven at 0.67 Hz. The nitrators were fitted with four-bladed propellers of 0.61 m diameter driven at 2 Hz. For exoling, the film coefficient h, for the inside of the vessel was given by:

$$\frac{h_b d_u}{k} \left(\frac{\mu_s}{\mu} \right)^{0.14} = 0.55 \left(\frac{L^2 N \rho}{\mu} \right)^{0.67} \left(\frac{C_\rho \mu}{k} \right)^{0.25} \tag{9.208}$$

which is very similar to that given by equation 9.207.

The film coefficients for the water jacket were in the range 635–1170 W/m² K-for water rates of 1.44–9.23 l/s. respectively. It may be noted that 7.58 l/s corresponds to a vertical velocity of only 0.061 m/s and to a Reynolds number in the annulus of 5350. The thermal resistance of the wall of the pan was important, since with the sulphonator it accounted for 13 per cent of the total resistance at 32.3 K and 31 per cent at 40.3 K. The change in viscosity with temperature is important when considering these processes, since, for example, the viscosity of the sulphonation liquors ranged from 340 mN s/m² at 32.3 K to 22 mN s/m² at 40.3 K.

In discussing equations 9,207 and 9,208 FLETCHER⁽⁹⁶⁾ has summarised correlations obtained for a wide range of impeller and agitator designs in terms of the constant before the Reynolds number and the index on the Reynolds number as shown in Table 9,11.

Table 9.11. Data on common agitators for use in equations 9.207

Type of agitator	Constant	Index
Flat blade disc turbine		
unhailled, or bailled vessel, Re < 4(8)	25.0	17.0
baffled, Re > 400	100	6.0%
Retreating-blade turbine with three	0.74	(O'D
blades, jacketed and haffled vessel.		
Re = 2 × 10° to 2 × 10°		
glassed steel impeller	0.33	200
affer other image in	6.00	0.0
anay seed impeller	0.37	69.0
Propeller with three blades		0
baffled vessel, Re = 5500 to 17 (vv)	140	
Plut blade paddle	1670	9.07
baffled or unbaffled yessel. Re > 2008)	7 V	8
Anchor	6.30	6.0
Re == 30 to 300	50	9
Re = 300 to 5000	8 e	0.00

9.8.3. Time required for heating or cooling

It is frequently necessary to heat or cool the contents of a large batch reactor or storage and. In this case the physical constants of the liquor may alter and the overall transfer coefficient may change during the process. In practice, it is often possible to assume an average value of the transfer coefficient so as to simplify the calculation of the time required for heating or cooling. The heating of the contents of a storage tank is cominonly effected by condensing steam, either in a coil or in some form of hairpin tube

The case of a storage tank with liquor of mass m and specific heat C_p , heated by seam condensing in a helical coil, it may be assumed that the overall transfer coefficient U is constant. If T_s is the temperature of the condensing steam, T_1 and T_2 the initial and final temperatures of the liquor, and A the area of heat transfer surface, and T_s the temperature of the liquor at any time t, then the rate of transfer of heat is given by:

$$Q = mC_p \frac{dT}{dt} = UA(T_s - T)$$

$$\frac{dT}{dt} = \frac{UA}{mC_p} (T_s - T)$$

$$\int_{T_1}^{T_2} \frac{dT}{T_s - T} = \frac{UA}{mC_p} \int_0^t dt$$

$$\ln \frac{T_s - T_1}{T_s - T_2} = \frac{UA}{mC_p} t$$
(9.2)

From this equation, the time t of heating from T_1 to T_2 , may be calculated. The same analysis may be used if the steam condenses in a jacket of a reaction vessel.

This analysis does not allow for any heat losses during the heating, or, for that matter, cooling operation. Obviously the higher the temperature of the contents of the vessel, the grader are the heat losses and, in the limit, the heat supplied to the vessel is equal to the fact losses, at which stage no further rise in the temperature of the contents of the vessel is possible. This situation is illustrated in Example 9.25.

The heating-up time can be reduced by improving the rate of heat transfer to the fluid. It agitating of the fluid for example, and by reducing heat losses from the vessel by misulation. In the case of a large vessel there is a limit to the degree of agitation possible, and circulation of the fluid through an external heat exchanger is an attractive alternative.

Example 9.25

A veset contains 1 tonne; (1 Mg) of a liquid of specific heat capacity 4.0 kJ/kg K. The vesset is heated by sean at 35.3 K which is fed to a cost innersed in the agitated liquid and heat is lost to the surroundings at 29.3 K from the outside of the vessel. How long does it take to hear the liquid from 293 to 35.3 K and what is the brakimum temperature to which the liquid can be heated? When the liquid temperature has reached 35.3 K, the steam supply is turned off for 2 hours (7.2 kg) and the vessel cools. How long will it take to reheat the instant at 35.3 K? The surface area of the coil is 0.5 m² and the overall exclicient of heat transfer to the liquid may be taken as 600 W/m² K. The outside area of the vessel is 6 m² and the exclinion of heat transfer to the liquid may be taken as 10 W/m² K.

Solution

If T K is the temperature of the liquid at time t s, then a heat balance on the vessel gives;

$$(1000 \times 4000) \frac{dT}{dt} = (600 \times 0.5)(393 - T) - (10 \times 6)(T - 293)$$

$$4,000,000 \frac{dT}{dt} = 135,480 - 360T$$

$$\frac{dT}{dt} = 376.3 - T.$$

The equilibrium temperature occurs when dT/dt = 0,

that is when:

In heating from 293 to 353 K, the time taken is:

$$t = 11.111 \int_{293}^{333} \frac{dT}{(376.3 - T)}$$
$$= 11.111 \ln \left(\frac{83.3}{23.3}\right)$$
$$= 14.155 \text{ (or 3.93 h)}.$$

The steam is turned off for 7200 s and during this time a heat balance gives:

$$(1000 \times 4000) \frac{dT}{dt} = -(10 \times 6)(T - 293)$$

 $66.700 \frac{dT}{dt} = 293 - T$

The change in temperature is then given by:

$$\int_{559}^{T} \frac{dT}{(293 - T)} = \frac{1}{66,700} \int_{0}^{7200} dt$$

$$\ln \frac{-60}{293 - T} = \frac{7200}{66,700} = 0.108$$

T = 346.9 K.The time taken to reheat the liquid to 353 K is then given by:

mg

$$t = 11.111 \int_{3450}^{335} \frac{dT}{(376.3 - T)}$$
$$= 11.111 \ln \left(\frac{29.4}{23.3}\right)$$
$$= 2584 \text{ s} \quad (0.72h).$$

9.9. SHELL AND TUBE HEAT EXCHANGERS

9.9.1. General description

Since shell and tube heat exchangers can be constructed with a very large heat transfera surface in a relatively small volume, fabricated from alloy steels to resist corrosion and

HEAT TRANSFER

be used for heating, cooling and for condensing a very wide range of fluids, they are the most widely used form of heat transfer equipment. Figures 9.62-9.64 show various forms of construction and a tube bundle is shown in Figure 9.65. The simplest type of unit shown in Figure 9.62, has fixed tube plates at each end into which the tubes are expanded. The tubes are connected so that the internal fluid makes several passes up and

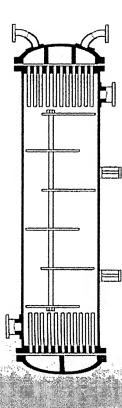


Figure 9.62. Heat exchanger with fixed tube plates (four tube, one shell-pass)

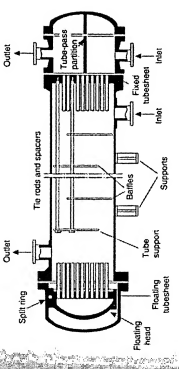


Figure 9.63. Heat exchanger with floating head (two tube-pass, one shell-pass)

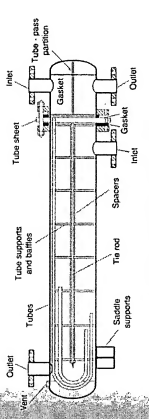


Figure 9.64. Heat exchanger with hairpin tubes

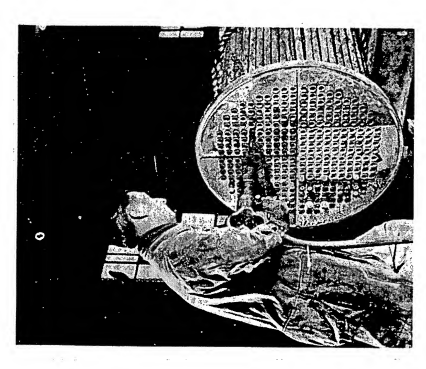


Figure 9.65. Expanding the ends of the tubes into the tube plate of a heat exchanger bundle

in one sense and then in the opposite sense across the tube bundle by fitting a series of baffles along the length. These baffles are frequently of the segmental form with about down the exchanger thus enabling a high velocity of flow to be obtained for a given heat transfer area and throughput of fluid. The fluid flowing in the shell is made to flow first 25 per cent cut away, as shown in Figure 9.29 to provide the free space to increase the velocity of flow across the tubes, thus giving higher rates of heat transfer. One problem with this type of construction is that the tube bundle cannot be removed for cleaning and no provision is made to allow for differential expansion between the tubes and the shell although an expansion joint may be fitted to the shell.

In order to allow for the removal of the tube bundle and for considerable expansion of one tube plate is fixed as before, but the second is bolted to a floating head cover so hat the tube bundle can move relative to the shell. This floating tube sheet is clamped the tubes, a floating head exchanger is used, as shown in Figure 9.63. In this arrangement

between the floating head and a split backing flange in such a way that it is relatively easy to break the flanges at both ends and to draw out the tube bundle. It may be noted that the shell cover at the floating head end is larger than that at the other end. This enables nember to be placed as near as possible to the edge of the fixed tube plate, leaving very

HEAT TRANSFER

as shown in Figure 9.64. This design is very commonly used for the reboilers on large Another arrangement which provides for expansion involves the use of hairpin tubes. little unised space between the outer ring of tubes and the shell.

in these designs there is one pass for the fluid on the shell-side and a number of passes on the tube-side. It is often an advantage to have two or more shell-side passes, although this considerably increases the difficulty of construction and, very often therefore, several factionating columns where steam is condensed inside the tubes.

Against this, the cost of pumping increases rapidly with increase in velocity and an son of a unit which is reliable and has the desired capacity, and secondly, the need to components and fittings and making the design as simple as possible. In most cases, it economic balance must be struck. A typical graph showing the operating costs, depreciation and the total cost plotted as a function of the water velocity in the tubes is shown arms cost. Thus in a condenser, for example, a high heat transfer coefficient is obtained and hence a small exchanger is required if a higher water velocity is used in the tubes. The essential requirements in the design of a heat exchanger are, firstly, the proviprovide an exchanger at minimum overall cost. In general, this involves using standard is necessary to balance the capital cost in terms of the depreciation against the opersmaller exchangers are connected together to obtain the same effect. in Figure 9.66.

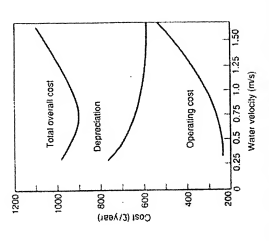


Figure 9.66. Effect of water velocity on unnual operating cost of condenser

HEAT TRANSFER

9.9.2. Basic components

The various components which make up a shell and tube heat exchanger are shown in of the record that noting that SAUNDERS⁽⁹³⁾ has presented a detailed discussion of Figures 9.63 and 9.64 and these are now considered. Many different mechanical arrange. ments are used and it is convenient to use a basis for classification. The standard published by the Tubular Exchanger Manufacturer's Association (TEMA(971) is outlined here!! design codes and problems in fabrication.

methods are based, although these may be adapted for other shell types by allowing Of the various shell types shown in Figure 9.67, the simplest, with entry and exit nozzie, at opposite ends of a single pass exchanger, is the TEMA E-type on which most design for the resulting velocity changes. The TEMA F-type has a longitudinal baffle giving two shell passes and this provides an alternative arrangement to the use of two shell required in order to cope with a close temperature approach or low shell-side flowrates. The pressure drop in two shells is some eight times greater than that encountered in the E-type design although any potential leakage between the longitudinal baffle and the shell in the F-type design may restrict the range of application. The so-called "split-flow" type of unit with a longitudinal baffle is classified as the TEMA G-type whose performance for reboilers and only occasionally for systems where there is no change of phase. The hence the shell-side fluid is in pure counterflow giving extremely low pressure drops and superior although the pressure drop is similar to the E-type. This design is used mainly with a pressure drop some one-eighth of the E-type, finds application in gas coolers and condensers operating at low pressures. The TEMA X-type shell has no cross haffles and so-called "divided-flow" type, the TEMA J-type, has one inlet and two outlet nozzles and again, this type of design is used for gas cooling and condensation at low pressures.

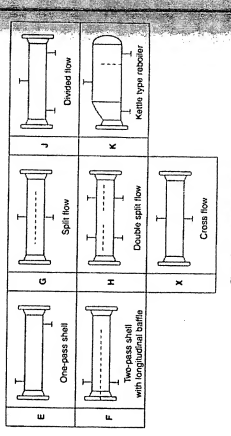


Figure 9.67. Teata shell types

The shell of a heat exchanger is commonly made of carbon steel and standard pipes are used for the smaller sizes and rolled welded plate for the larger sizes (say 0.4-1.0m)

The hickness of the shell may be calculated from the formula for thin-walled cylinders and a minimum thickness of 9.5 mm is used for shells over 0.33 m o.d. and 11.1 mm for shells over 0.9 m o.d. Unless the shell is designed to operate at very high pressures, the enculated wall thickness is usually less than these values although a corrosion allowance of 32 mm is commonly added to all carbon steel parts and thickness is determined more bengidity requirements than simply internal pressure. The minimum shell thickness for various materials is given in BS3274(99). A shell diameter should be such as to give as close a fit to the tube bundle as practical in order to reduce bypassing round the outside of the bundle. Typical values for the clearance between the outer tubes in the bundle and the inside diameter of the shell are given in Figure 9.68 for various types of exchanger.

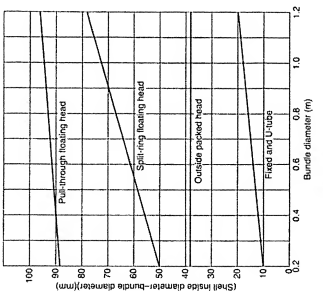


figure 9.68. Shell-bundle clearance

and the shell which cannot be tolerated where high purity or uncontaminated materials are required. In general, tube bundles make use of a fixed tubesheet, a floating-head or side pressures since these will both affect any potential leakage between the tube bundle Utubes which are shown in Figures 9.62, 9.63 and 9.64 respectively. It may be noted fere that the thickness of the fixed tubesheet may be obtained from a relationship of The detailed design of the tube bundle must take into account both shell-side and tubethe form:

$$d_1 = d_G \sqrt{0.25P/f}$$
 (9)

where d_G is the diameter of the gasket (m), P the design pressure (MN/m²). f the bottom of the partition plate grooves. The thickness of the floating head tubesheet is very allowable working stress (MN/m²) and d_i the thickness of the sheet measured at often calculated as $\sqrt{2d_i}$.

In selecting a tube diameter, it may be noted that smaller tubes give a larger heat transfer $6 \, \mathrm{mm} \, (\frac{5}{8} \, \mathrm{in})$ to 50 mm (2 in) O.D.; the smaller diameter usually being preferred as these increases the likelihood of tube vibration. Heat exchanger tubes are usually in the range. fluids. The tube thickness or gauge must be such as to withstand the internal pressure. area for a given shell, although 19 mm o.d. tubes are normally the minimum size used in order to permit adequate cleaning. Although smaller diameters lead to shorter tubes more holes have to be drilled in the tubesheet which adds to the cost of construction and clean especially by mechanical methods and are therefore widely used for heavily fouling and also to provide an adequate corrosion allowance. Details of steel tubes used in hear give more compact and therefore cheaper units. Against this, larger tubes are easied exchangers are given in BS3606(190) and summarised in Table 9.12, and standards other materials are given in BS3274(99).

Table 9.12. Standard dimensions of steel tubes

() ulstele districted	Bameler			Carrier sea	Street, and an area		45
do		Wall th	Wall thickness	100 Kg	Cross sectional area for flow	Surface unit	Surface are per unit length
(mm)	(in)	(mm)	(in)	(m ²)	(iii.)	(m,,/m)	(11/3)
91	0.630	77	0.047	0.000145	0.00156	0.0503	0.165
		9.1	0.063	0.000129	0.00139		
		2.0	0.079	0.000113	0.00122		A fig.
20	0.787	9.1	0.063	0.000222	0.00239	0.0628	2000
		2.0	0.079	0.000201	0.00216		
		2.6	0.102	0.000172	0.00185		
25	0.984	9.1	0.063	0.000373	0.00402	0.0785	926 0
		2,0	0.079	0.000346	0.00373		
		2.6	0.102	0.000308	0.00331		
		3.2	0.126	0.000272	0.00293		900
æ	<u>=</u>	9.	0.063	0.000564	0.00607	0.0942	DUL O
		2.0	0.079	0.000531	0.00572		
		2.6	0.102	0.000483	0.00512		* en
;		3.2	0.126	0.000437	0.00470		
æ.	1.496	5.0	0.079	0.000908	0.00977	0.1194	107.0
		2.6	0.102	0.000845	0160000		
		3.2	0.126	0.000784	0.00843		1000
50	1.969	2.0	0.079	0.001662	6821070	0.1571	5150
		2.6	0.102	0.001576	0.01697		
		3.2	0.126	0.001493	0.01607		

surface area due to the smaller shell diameter, the thinner tube sheets and flanges and the smaller number of holes to be drilled, and the reduced complexity. Preferred tube lengths are 1.83 m (6 ft), 2.44 m (8 ft), 3.88 m (12 ft) and 4.88 m (16 ft); larger sizes are used where the total tube-side flow is low and fewer, longer tubes are required in order In general, the larger the tube length, the lower is the cost of an exchanger for a given to obtain a required velocity. With the number of tubes per tube-side pass fixed in order to obtain a required velocity, the total length of tubes per tube-side pass is determined by the heat transfer surface required. It is then necessary to fit the tubes into a suitable shell.

v,

give the desired shell-side velocity. It may be noted that with long tube lengths and adjainely few tubes in a shell, it may be difficult to arrange sufficient baffles for adequate support of the tubes. For good all-round performance, the ratio of tube length to shell ameter is usually in the range 5-10.

HEAT THANSFER

minimum web thickness between tubes of about 3.2 mm to ensure adequate strength for the rolling. In general, the smallest pitch in triangular 30° layout is used for clean fluids In both laminar and turbulent flow and a 90° or 45° layout with a 6.4 mm clearance where mechanical cleaning is required. The bundle diameter, d_b , may be estimated from the equilateral triangular, square and staggered square arrays. The triangular layout gives a esign, the square array simplifies maintenance and particularly cleaning on the shellside. Good practice requires a minimum pitch of 1.25 times the tube diameter und/or a moust tube sheet although, because the vertical and horizontal distances between adjacent ubes is generally greater in a square layout compared with the equivalent triangular pitch Signbe layout and pitch, considered in Section 9.4.4 and shown in Figure 9.69, make use following empirical equation which is based on standard tube layouts:

Number of tubes,
$$N_t = a(d_b/d_o)^b$$
 (9.21)

where the values of the constants a and b are given in Table 9.13. Tables giving the number of tubes that can be accommodated in standard shells using various tube sizes. gitches and numbers of passes for different exchanger types are given, for example, in ERNCS and LUDWIG(101).

Number of passes	,	,		,
	4	4	c	x
Triangular pitch* a 0.319	0.249	0.175	0.0743	0.0365
b 2.142	2,207	2,285	2.499	2.675
Square pitch* a 0.215	0.156	0.158	0.0402	0.0331
P	2.291	2.263	1.617	2,643

Various buffle designs are shown in Figure 9.70. The cross-baffle is designed to direct the flow of the shell-side fluid across the tube bundle and to support the tubes against agging and possible vibration, and the most common type is the segmental baffle which provides a baffle window. The ratio, baffle spacing/baffle cut, is very important in maximising the ratio of heat transfer rate to pressure drop. Where very low pressure ipps are required, double segmental or "disc and doughnut" baffles are used to reduce the pressure drop by some 60 per cent. Triple segmental baffles and designs in which all the tubes are supported by all the baffles provide for low pressure drops and minimum tube vibration. * 2

With regard to buffle spacing. TEMA197 recommends that segmental baffles should not espaced closer than 20 per cent of the shell inside diameter or 50 mm whichever is the greater and that the maximum spacing should be such that the unsupported tube lengths, given in Table 9.14, are not exceeded. It may be noted that the majority of failures due to vibration occur when the unsupported tube length is in excess of 80 per cent of the FIEMA maximum; the best solution is to avoid having tubes in the baffle window.

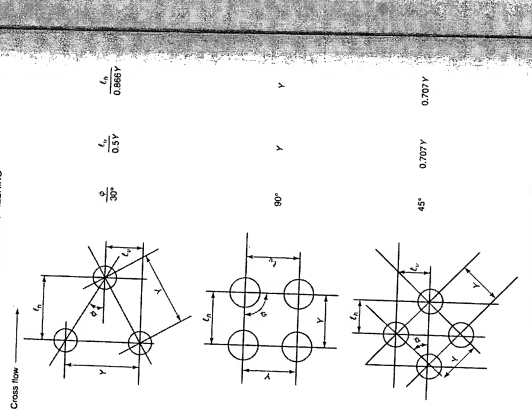
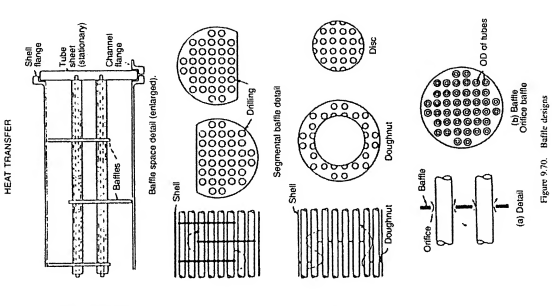


Figure 9.69. Examples of tube arrays(97)

9.9.3. Mean temperature difference in multipass exchangers

In an exchanger with one shell pass and several tube-side passes, the fluids in the tubes and shell will flow co-currently in some of the passes and countercurrently in the others. For given inlet and outlet temperatures, the mean temperature difference for countercurrent flow is greater than that for co-current or parallel flow, and there is no easy way of



Inding the true temperature difference for the unit. The problem has been investigated by UNDERWOOD⁽¹⁰²⁾ and by BOWMAN et al.⁽¹⁰³⁾ who have presented graphical methods for calculating the true mean temperature difference in terms of the value of θ_m which would be obtained for countercurrent flow, and a correction factor F. Provided the following conditions are maintained or assumed, F can be found from the curves shown in Figures 9.71–9.74.

Table 9.14. Maximum unsupported spans for tubes

Group A: Carbon and high alloy steel, low alloy steel, nickel-Materials

cepper, nickel, nickel-chronium-iron. Group B: Aluminium and aluminium alloys, copper and copper alloys, titanium and zirconium,

0,

0.9

0.8

Correction factor (E)

0.

9.0

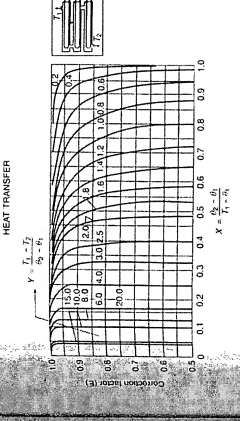


figure 9.73. Correction for logarithmic mean temperature difference for three shell pass exchanger

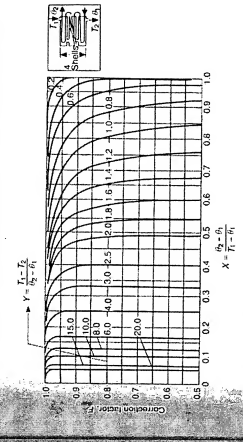


Figure 9.71. Correction for logarithmic mean temperature difference for single shell pass exchanger.

6.0

0.8

Correction factor, F

0.9

0.8

0,7

9.0

0,4

0.3

0

0.5

 $X = \frac{1}{T_1 - \theta_1}$ 020. 0.5

0.2

Figure 9.74. Correction for logarithmic mean temperature difference for four shell pass exchanger

(a) The shell fluid temperature is uniform over the cross-section considered as constituting a pass.

(b) There is equal heat transfer surface in each pass.

0.

0.0

0.8

0.7

9.0

0.5 0.4

S 0.2

0

0.5

0.6

0.7

(c) The overall heat transfer coefficient U is constant throughout the exchanger.

(d) The heat capacities of the two fluids are constant over the temperature range.

(c) There is no change in phase of either fluid. (f) Heat losses from the unit are negligible.

Figure 9.72. Carrection for logarithmic mean temperature difference for double shell pass exchanger.

Q = UAFO,

(9.212)

F is expressed as a function of two parameters:

$$X = \frac{\theta_2 - \theta_1}{T_1 - \theta_1}$$
 and $Y = \frac{T_1 - T_2}{\theta_2 - \theta_1}$ (9.2)

the temperature profile is as shown. Because one of the passes constitutes a parallel flow fluid temperature T₁. This is true for the conditions shown in Figures 9.75a and 9.75b arrangement, the exit temperature of the cold fluid 0, cannot closely approach the ha for three shell-side passes Figure 9.73, and for four shell-passes Figure 9.74. For the cas If a one shell-side system is used Figure 9.71 applies, for two shell-side passes Figure 9.7 of a single shell-side pass and two tube-side passes illustrated in Figures 9.75a and 9. and UNDERWOOD⁽¹⁰²⁾ has shown that F is the same in both cases.

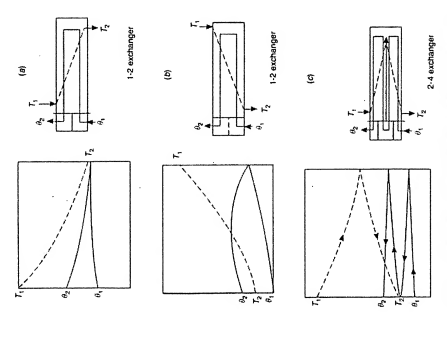


Figure 9.75. Temperature profiles in single and double shell pass exchangers

for example, an exchanger is required to operate over the following temperatures:

HEAT TRANSFER

$$T_1 = 455 \text{ K}$$
, $T_2 = 372 \text{ K}$

$$\theta_1 = 283 \text{ K}, \ \theta_2 = 388 \text{ K}$$

$$X = \frac{\theta_2 - \theta_1}{T_1 - \theta_1} = \frac{388 - 283}{455 - 283} = 0.6$$

$$V = T_1 - T_2 = 455 - 372 = 0$$

For a single shell pass arrangement, from Figure 9.71 F is 0.65 and, for a double shell $= \frac{388 - 283}{388 - 283} = 0.8$ $\theta = \theta_1$ 7 || -

by taking as an example the following data where equal ranges of temperature are considered: In order to obtain maximum heat recovery from the hot fluid, θ_2 should be as high as ossible. The difference $(T_2 - \theta_2)$ is known as the approach temperature and, if $\theta_2 > T_2$, apidly when there is but a single pass on the shell-side. This implies that, in parts of the heat exchanger, heat is actually being transferred in the wrong direction. This may be illustrated ass arrangement, from Figure 9.72 F is 0.95. On this basis, a two shell-pass design then a temperature cross is said to occur; a situation where the value of F decreases very

Case, T	T_2	6	θ_2	Approach	×	٨	F
- 531 - 42 - 23				$(T_2-\theta_2)$			
19 1	513	363	463	.50	0.4		0.92
573	473	373	473	. 0	0.5		0.80
3 (1) 543	443	363	463	cross of 20	0.55	-	0.66

passes on the shell-side are required. On the very largest installations it may be necessary to inking a number of exchangers in parallel arranged as, say, sets of three in series as shown in passes should be used. It is seen from Figure 9.75b that there may be some point where the emperature of the cold fluid is greater than θ_2 so that beyond this point the stream will be gooled rather than heated. This situation may be avoided by increasing the number of shell passes. The general form of the temperature profile for a two shell-side unit is as shown in figure 9.75c. Longitudinal shell-side baffles are rather difficult to fit and there is a serious hance of leakage. For this reason, the use of two exchangers arranged in series, one below the other, is to be preferred. It is even more important to employ separate exchangers when three Tigure 9.76. This arrangement is preferable for any very large unit which would be unwieldy as a single system. When the total surface area is much greater than 250 m², consideration If a temperature cross occurs with a single pass on the shell-side, a unit with two shell should be given to using multiple smaller units even though the initial cost may be higher.

network so as to reduce to a minimum both the heating and cooling duties placed on the will be some point where the temperature difference between the hot and cold streams is a intotal, there is a net surplus or deficit of heat available. It is of great economic importance Dischieve the most effective match of the hot and cold streams in the heat exchanger works utilities, such as supplies of steam and cooling water. This necessitates making the best use of the temperature driving forces. In considering the overall requirements there In many processing operations there may be a large number of process streams, some of Which need to be heated and others cooled. An overall heat balance will indicate whether,

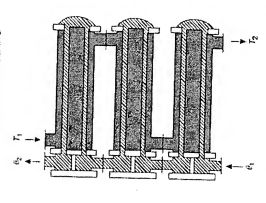


Figure 9.76. Set of three heat exchangers in series

minimum and this is referred to as the *pinch*. The lower the temperature difference autapinch point, the lower will be the demand on the utilities, although it must be remembered that a greater area, (and hence cost) will be involved and an economic balance-must therefore be struck. Heat exchanger networks are discussed in Volume 6 and, in detain the User Guide published by the Institution of Chemical Engineers⁽¹⁰⁴⁾. Subsequently, LINNHOFF⁽¹⁰⁵⁾ has given an overview of the industrial application of pinch analysis to the design of networks in order to reduce both capital costs and energy requirements.

Example 9.26

Using the data of Example 9.1, calculate the surface area required to effect the given duty using a multipact heat exchanger in which the cold water makes two passes through the tubes and the hot water makes assure pass through the shell.

Solution

As in Example 9.1, the heat load = 1672 kW

With reference to Figure 9.71: $T_1 = 360 \text{ K}$, $T_2 = 340 \text{ K}$

$$X = \frac{\theta_2 - \theta_1}{T_1 - \theta_1} = \frac{316 - 300}{360 - 300} = 0.267$$

and hence:

$$Y = \frac{T_1 - T_2}{\theta_2 - \theta_1} = \frac{360 - 340}{316 - 300} = 1.25$$

and:

from Figure 9.58:

F = 0.97

HEAT TRANSFER

 $F\theta_m = (41.9 \times 0.97) = 40.6 \text{ K}$

1672

 $A = \frac{10.12}{2 \times 40.6} = \frac{20.6 \text{ m}^2}{20.00}$

the beat transfer area is then:

9:9:4. Film coefficients

practical values

ranser coefficient U and the correction factor F in equation 9.212. The determination mo methods have been developed for measuring film coefficients. The first requires a the temperatures of both fluids and the surface of separation. With a concentric tube be noted that when the heat flux is very high, as with boiling liquids, there will be un appreciable temperature drop across the tube wall and the position of the thermocouple is in any firm of heat transfer equipment, the required area of heat transfer surface for given load is determined by the overall temperature difference θ_n , the overall hear of the individual film coefficients which determine the value of U has proved difficult even for simple cases, and it is quite common for equipment to be designed on the hass of practical values of U rather than from a series of film coefficients. For the important case of the transfer of heat from one fluid to another across a metal surface. mowledge of the temperature difference across each film and therefore involves measuring system it is very difficult to insert a thermocouple into the thin tube and to prevent the thermocouple connections from interfering with the flow of the fluid. Nevertheless, this method is commonly adopted, particularly when electrical heating is used, It must then important. For this reason, working with stainless steel, which has a relatively low themal conductivity, is difficult.

The second method uses a technique proposed by WiLSON(106). If steam is condensing on the guiside of a horizontal tube through which water is passed at various velocities, then the overall and film transfer coefficients one related by:

$$\frac{1}{U} = \frac{1}{h_o} + \frac{x_w}{k_w} + R_i + \frac{1}{h_i}$$
 (from equation 9.201)

provided that the transfer area on each side of the tube is approximately the same.

For conditions of turbulent flow the transfer coefficient for the water side, $h_i = \varepsilon u^{0.8}$, R_i the scale resistance is constant, and h_o the coefficient for the condensate film is almost independent of the water velocity. Thus, equation 9.201 reduces to:

$$\frac{1}{U} = (\text{constant}) + \frac{1}{\epsilon u^{0.8}}$$

If $I(U_i)$ plotted against $1/u^{0.8}$ a straight line, known as a Wilson plot, is obtained with slope of $1/\epsilon$ and an intercept equal to the value of the constant. For a clean tube R_i should be nil, and hence I_0 , can be found from the value of the intercept, as x_e/k_w will generally be small for a metal tube, h_i may also be obtained at a given velocity from the difference between 1/U at that velocity and the intercept.

This technique has been applied by RHODES and YOUNGER(107) to obtain the values of h. lor-condensation of a number of organic vapours, by PRATT⁽⁹²⁾ to obtain the inside coefficient for coiled tubes, and by COULSON and MEHTA⁽¹⁰⁸⁾ to obtain the coefficient for

1		١	<u>ت</u>	.		-			A. ee	
	Aluminium		0.0034	0.0074	0.0173	2	O Omnos.	0.0000	0.000053	0.000070
K/k/W)	Admiralty	metal	0.011	0.019	0.046	'F/Btu)	0.000065	0.000086	0.00011	07/10/170
Values of x, /k, (m2K/kW	Stainless		0.083	0.141	0.176	es of xw/kw (ft ² h°	0.00047	0.00062	0.000	
I ₂ V	Steel	0100	0.025	0.032	250.0	Value	0.0001	0.00014 8.00014	0.00024	
Ę	Copper	0.0031	0.0042	0.0055	* 10000	O CONTROL	0.000018	0.000031	0.000041	***************************************
Thickness	î mu	1.24	1.65	2.77		6700	0.065	0.083	0.109	
Gauge (BWG)		<u>so</u> ;	₽ <u>-</u> 2	12		<u>×</u>	9:	T :	77	

	III NXW	fr-h"F/Btu			
Water*				m*K/kW	fr'h"F/Bm
distilled	60.0	0.0005	Steam good quality, oil-		Pagi
clear river unfreated cooling	0.21	0.0012	free poor quality, oil-	0.052	0.0003
tower treated cooling	0.58	0.0033	free exhaust from	60.0	0.0005
tower treated boiler feed hard well	0.26 0.26	0.0015	reciprocating engines	0.18	
	0.38	0.0033	Liquids		
Gasses air	300		treated brine	0.27	0.0015
solvent vapours	0.14	0.0015-0.003	fuel ails	81.0	0.00 0.000

an annulus. If the results are repeated over a period of time, the increase in the value of R_i can also be obtained by this method

Typical values of thermal resistances and individual and overall heat transfer coefficients are given in Tables 9.15–9.18.

Correlated data

Heat transfer data for turbulent flow inside conduits of uniform cross-section are usually correlated by a form of equation 9.66;

$$Nu = CRe^{0.3} p_1^{0.33} (\mu/\mu_s)^{0.14}$$
 (9.21a)

where, based on the work of SIEDER and TATE⁽¹⁷⁾, the index for the viscosity correction term is usually 0.14 although higher values have been reported. Using values of C. of 0.021 for gases, 0.023 for non-viscous liquids and 0.027 for viscous liquids, equation 9.214 is sufficiently accurate for design purposes, and any errors are far outweighed by uncertainties in predicting shell-side coefficients. Rather more accurate tube-side data

HEAT TRANSFER

Table 9.17. Approximate overall heat transfer coefficients U for shell and tube equipment

Continuers Committee of grants colvents (atmospheric) Steam (vacuum) Steam (vacuum) Steam (vacuum) Organic solvents (atmospheric) Steam (organic solvents (atmospheric) Organic solvents (atmospheric and high Organic solvents (atmospheric and high Organic solvents (atmospheric and high Organic solvents (atmospheric) Organic solvents (atmospheric) Mater - hrine Organic solvents Steam		350-750 300-600 100-200 300-600 100-200 10-50 80-200 10-30 10-80 1
pheric) In some I high pheric) m)	_	350 - 750 300 - 600 100 - 200 50 - 120 20 - 80 10 - 50 80 - 200 10 - 30 10 - 80 10 - 8
m some m some I high th m)	_	330 – 750 300 – 600 100 – 200 30 – 120 20 – 80 10 – 30 10 – 80 10 – 30 10 – 80 10 – 200 10 –
pheric) m some l high ft m) m)		300-730 300-730 300-730 100-200 10-3
m some I high the control of the co	-	300-600 100-200 30-120 20-80 10-50 80-200 10-30 50-130 100-200 5-50 5-50 5-40
phenci) In some (h phenci) m) m)		20-20 20-120 20-80 10-30 80-200 10-30 10-30 10-200 5-50 5-50 5-50 5-50 5-50 5-50 5-50
m some thigh m) m)	-	20-120 20-80 10-50 80-200 10-30 10-30 100-200 5-50 5-50
th igh pheric) m)	_	\$0-120 20-80 10-50 80-200 10-30 50-150 100-200 5-50 4-40
Thigh pheric) m)		20-80 10-50 80-200 10-30 10-30 50-150 100-200 5-50 4-40
m) m)		20-80 10-50 80-200 10-30 10-30 50-150 100-200 5-50 5-50
m) pheric)		250-30 10-30 80-200 10-30 10-30 50-150 100-200 5-50 4-40
m)	-	10-50 80-200 10-30 10-30 59-150 100-200 5-50 5-50
m)		10-50 80-200 10-30 10-30 50-150 100-200 5-50 4-40
m) (ii)	<u> </u>	80-200 10-30 250-750 50-150 10-80 100-200 5-50 5-40
· ·		250-750 50-150 10-80 100-200 5-50 4-40
	_	259-750 50-150 10-80 100-200 5-50
	_	250-750 50-150 10-80 100-200 5-50 4-40
		250-730 50-150 100-200 5-50 5-50
		50-150 10-80 100-200 5-50 4-40
		100-20 100-20 5-50
		5-50 5-50 4-40
	30-300	5-50
	20-200	9440
	UUV = US	
	000	(K) = 0
	2000-4000	350-750
	8 600-1200	100-200
	400-1000	80-180
	uum) 150~400	25-75
	400000	75-150
	200-600	30-100
solvents ths oils	00071-0000	150. 300
sils oils	005-005	051-05
pils oils	005-00	S-1-20
	000-00	071 07
	001 07	(S) (S)
	00-200	06-01
	004-001	20-70
	600-1200	100-300
c solvents	200-500	30-90
		3-50
Organic solvents Organic solvents	100-400	20-60
Heavy oils	50-300	8-50

may be obtained by using correlations given by the Engineering Sciences Data Unit and, based on this work, BUTTERWORTH(109) offers the equation:

$$St = E \text{ Re}^{-0.205} p_r^{-0.505}$$
 (9.215)

where:

the Stanton Number $S_I = NuRe^{-1}Pr^{-1}$

 $E = 0.22 \exp[-0.0225(\ln Pr)^2]$

Equation 9.215 is valid for Reynolds Numbers in excess of 10,000. Where the Reynolds Number is less than 2000, the flow will be laminar and, provided natural convection effects

CHEMICAL ENGINEERING

Table 9.18. Approximate film coefficients for heat transfer

We change of state W/m² K Buuft²h ³F water Water 300-2000 gases 20-300 300-2000 oils 30-300 60-500 Goods 30-2000 60-500 steam 60-700 10-120 steam 6000-1700 100-300 organic solvents 900-2800 100-300 light oils 120-300 20-500 heavy oils (vacuum) 120-300 20-50 water 3000-1000 30-200 water 2000-12,000 100-300 organic solvents 600-2000 100-300 water 2000-1000 100-300 organic solvents 600-2000 100-300 light oils 800-1000 100-300 heavy oils 600-2000 100-300		h, 0	h, or ho
olvents 1700–11,000 20–300 20–300 20–300 20–300 60–700 20–300 2000–12,000 120–2300	***************************************	W/m² K	Btu/ft²h *F
1700 - 11,000 20 - 300 20 - 300 20 - 300 350 - 2000 60 - 700 10 350 - 2000 10 350 - 2000 10 350 - 2000 10 350 - 2300 120 - 2300 120 - 2300 120 - 2300 100 - 2300 100 - 1200 100	No change of state		
olvents 1700. 1700	Water	3.770	
olvents 270–300 on 60–700 on 600–2000 olvents 900–2800 olvents 900–2800 olvents 900–2800 olvents 600–2000 olvents	gases	000711-0071	300-2000
200-3000 m 600-700 slvenis 6000-17,000 slvenis 1200-300 120-300 3000-6000 2000-12,000 1100-2300 800-1760 60-300	Organic solvents	(06,-07	3.50
60–700) 600–700) 6000–17,000 600–2800 1200–2800 1200–300 1000–2000 1100–2300 800–1300 600–2000 600–300	Oils	350~3000	ODS09
6000-17,000 3lvenis 600,0-17,000 120-2300 120-300 3000-6000 3000-12,000 400-2300 1100-2300 8100-1700 601-3700 601-3700		60-700	10-120
Sivents 6000—17,000 1200—2300 1200—2300 120—300 3000—6000 2000—12,000 1100—2300 800—1200 601–1200 601–1200 601–1200	Condensation		071
100 100	steam	, CONTA	
(vacuum) 120300 120300 120300 30006000 1002000 11002300 8001200 600-300 90130	Organic solvente	CHAND-17 (NEE)	1000 3000
120-300 120-300 120-300 3000-6000 100-200 100-2300 100-130 100-130 100-130 100-130	light oils	WW-28(0)	150-500
120-300 3000-6000 (vents 600-2000 1100-2300 200-1700 1300-1700 100-300	heavy oils organis	1200-2300	200-400
3000~6000 2000~12,000 600~2000 1100~2300 800~1700 600~300	American (Vacuum)	120-300	30.50
1000 - 10	Total Maria	3000-6000	00-0-
2008-12,000 600-2000 1100-2300 800-1700 601-300	Vaporation	TATALAN A	200-1000
4vents 600–2000 100–2000 1100–2300 800–1700 60–300	Water	Tribo	
600-2300 100-2300 800-1700 60-1700	Organic sulvente	2000 - 12-000)	30-200
1100-2300 800-1700 60-300	ammonia	CHX)-2000	1001-300
800-1700 60:-100	Dehroits	1100-2300	200-400
90£~(K)	herve oils	S(N)-17(N)	150-200
	Sim francis	100 × (N)	7876

are negligible, film coefficients may be estimated from a form of equation 9.85 meding to take account of the variation of viscosity over the cross-section;

$$Nu = 1.86(RePr)^{0.33}(d/I)^{0.33}(\mu/\mu_s)^{0.14}$$

The minimum value of the Nusselt Number for which equation 9.216 applies is 33

Reynolds Numbers in the range 2000-10,000 should be avoided in designing hear exchangers as the flow is then unstable and coefficients cannot be predicted with any degree of accuracy. If this cannot be avoided, the lesser of the values predicted by

As discussed in Section 9.4.3, heat transfer data are conveniently correlated in terms of a heat transfer factor ja, again modified by the viscosity correction factor;

$$j_h = SIPr^{0.67}(\mu/\mu_s)^{-0.14}$$

which enables data for laminar and turbulent flow to be included on the same plot as to estimate coefficients with heat exchanger tubes and commercial pipes although, due to shown in Figure 9.77. Data from Figure 9.77 may be used together with equation 9.217. a higher roughness, the values for commercial pipes will be conservative. Equation 9.217 is rather more conveniently expressed as:

$$Nu = (hd/k) = j_h ReP r^{0.33}(\mu/\mu_s)^{-0.14}$$

It may be noted that whilst Figure 9.77 is similar to Figure 9.24, the values of jh differ.

 $i_H = NuPr^{-0.33}(\mu/\mu_3)^{-0.14}$ Thus the relationship between j_h and j_H is:

(9.220)

HEAT TRANSFER

discussed in Section 9.4.3, by incorporating physical properties into equations 9.214 2016, correlations have been developed specifically for water and equation 9.221, and data from EAGLE and FERGUSON⁽¹¹⁰⁾ may be used:

$$h = 4280(0.00488T - 1)n^{0.3}/d^{0.2}$$
 (9.22)

when is in SI units, with h (film coefficient) in W/m²K, T in K, u in ms and d in m.

Estimate the heat transfer area required for the system considered in Examples 9.1 and 9.26, assuming that no fact the overall coefficient of heat transfer are available.

As in the previous examples.

heat lond = 1672 kW

corrected mean temperature difference. $F\theta_{\rm m}=40.6~{\rm deg~K}$

mean water temperature, $T = 0.5(360 \pm 340) = 350 \text{ K}$

ing a tube diameter, d = 19 mm or 0.00119 m and a water velocity, u = 1 m/s, then, in equation 9.221;

 $h_{\rm i} = 4280 (0.00 \pm 88 \times 350) - 11.0^{6.8} / 0.0019^{0.2} = 10610 \text{ W/m}^2 \text{K or } 10.6 \text{ kW/m}^2 \text{K}$

From Table 9.18, an estimate of the shell-side film coefficient is:

$$h_0 = 0.501700 + 11000) = 6353 \text{ W/m}^2\text{K} \text{ or } 6.35 \text{ kW/m}^2\text{K}$$

Fristed tubes of a wall thickness of 1.6 mm, the thermal resistance of the wall, from Table 9.15 is:

$$x_w/k_w = 0.025 \text{ m}^2 \text{KAW}$$

and the thermal resistance for treated water, from Table 9.16, is 0.26 m²KkW for both layers of scale. Thus, in Equation 9.201:

 $(1/U) = (1/h_0) + (x_u/k_u) + R_i + R_0 + (1/h_i)$

$$= (1/6.35) + 0.025 + 0.52 + (1/10.6) = 0.797 \text{ m}^2\text{K/kW}$$

 $U = 1.25 \text{ kW/m}^2$ The heat transfer area required is then:

 $U = 1.25 \text{ kW/m}^2 \text{K}$

$$A = Q/F\theta_m U = 1672/(40.6 \times 1.25) = 32.9 \text{ m}^2$$

As discussed in Section 9.4.4, the complex flow pattern on the shell-side and the great Jumber of variables involved make the prediction of coefficients and pressure drop very difficult, especially if leakage and bypass streams are taken into account. Until about 1960, empirical methods were used to account for the difference in the performance

of real exchangers as compared with that for cross-flow over ideal tube banks. The nethods of Kern⁽¹³⁾ and Donortue⁽¹¹⁾ are typical of these "bulk flow" methods and dear approach, together with more recent methods involving an analysis of the contribution to heat transfer by individual streams in the shell, are discussed in Section 9.9.6.

Special correlations have also been developed for liquid metals, used in recent years in the nuclear industry with the aim of reducing the volume of fluid in the heat transfer grants. Such fluids have high thermal conductivities, though in terms of heat capacity per anti-volume, liquid sodium, for example, which finds relatively widespread application, has a value of $C_p\rho$ of only 1275 kJ/m^3 K.

Although water has a much greater value, it is unsuitable because of its high vapour pressure at the desired temperatures and the corresponding need to use high-pressure piping. Because of their high thermal conductivities, liquid metals have particularly low values of the Prandtl number (about 0.01) and they behave rather differently from normal nuits, under conditions of forced convection. Some values for typical liquid metals are given in Table 9.19.

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Table 9.19. Prandtl numbers of liquid metals

:	Temperature	Prundtt number
Metal	(K)	Pr
Potassium	975	0.003
Sectium	975	0000
Na/K altoy (56:44)	975	900
Mercury	57.5	0.008
Lithium	475	0.065

The results of work on sodium, lithium, and mercury for forced convection in a pipe have been correlated by the expression:

)

$$Nu = 0.625(RePr)^{0.4} (9.222$$

although the accuracy of the correlation is not very good. With values of Reynolds number of about 18,000 it is quite possible to obtain a value of h of about 11 kW/m² K for flow in a pipe:

9.9.5. Pressure drop in heat exchangers

Tube-side

Pressure drop on the tube-side of a shell and tube exchanger is made up of the friction loss in the tubes and losses due to sudden contractions and expansions and flow reversals experienced by the tube-side fluid. The friction loss may be estimated by the methods outlined in Section 3.4.3 from which the basic equation for isothermal flow is given by equation 3.18 which can be written as:

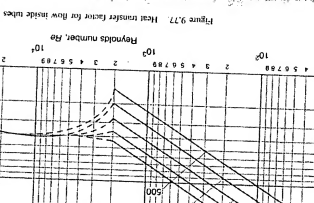
$$-\Delta P_t = 4 j_f(l/d_i)(\rho u^2)$$
 (9.223)

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Heat transfer factor, j_h

10.3

where j_f is the dimensionless friction factor. Clearly the flow is not isothernal and it is usual to incorporate an empirical correction factor to allow for the change in physical



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CHEMICAL ENGINEERING

524

properties, particularly viscosity, with temperature to give:

$$-\Delta P_i = 4j_f (I/d_i)(\rho u^2)(\mu/\mu_i)^m$$

2100). Values of j_f for heat exchanger tubes are given in Figure 9.78 which is based where m=-0.25 for laminar flow (Re < 2100) and -0.14 for turbulent flow.

velocity heads. In this way, the total loss is 10,5 velocity heads, or 2.6 per pass, giving There is no entirely satisfactory method for estimating losses due to contraction and given in Table 3.2. With four tube passes, for example, there will be four contrating BUTTERWORTH [13] 1.8. LORD et al. [114] suggests that the loss per pass is equivalent adding four velocity heads per pass, FRANK⁽¹¹²⁾ recommends 2.5 velocity heads a tube length of 300 diameters for straight tubes and 200 for U-tubes, whilst Evays recommends the addition of 67 tube diameters per pass. Another approach is to estimate equivalent to a loss of $(4 \times 0.5) = 2$ velocity heads, four expansions equivalent to a loss the number of velocity heads by using factors for pipe-fittings as discussed in Section the tube inlets, expansion at the exits and flow reversals, although KERN⁽²⁸⁾ sugg $(4 \times 1.0) = 4$ velocity heads and three 180° bends equivalent to a loss of (3×1.5) support to Frank's proposal of 2.5. Using this approach, equation 9.224 becomes:

$$-\Delta P_{\text{total}} = N_P[4j_f(l/d_i)(\mu/\mu_s)^m + 1.25](\rho u^2)$$

(9.23) where NP is the number of tube-side passes. Additionally, there will be expansion in estimated by adding one velocity head for the inter, and 0.5 of a velocity head for the outlet, based on the nozzle velocities. Losses in the mount contraction losses at the inlet and outlet nozzles respectively, and these losses may outlet, based on the nozzle velocities. Losses in the nozzles are only significant for ga

Shell-side

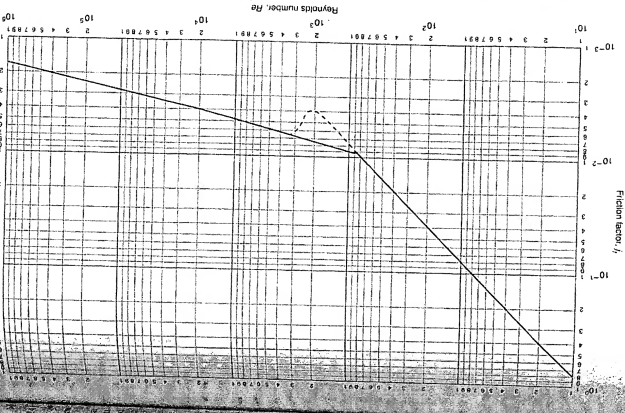
As discussed in Section 9.4.4, the prediction of pressure drop, and indeed heat transfer in the segmentally baffled unit. Whilst the baffles are intended to direct fluid across the tubes, the actual flow is a combination of cross-flow between the baffles and axial or parallel flow in the baffle windows as shown in Figure 9.79, although even this does in Figure 9.80 which is based on the work of Tinker (116) who identifies the various stream. coefficients, in the shell is very difficult due to the complex nature of the flow patien the fabrication and assembly of the unit. This more realistic flow pattern is shown represent the actual flow pattern because of leakage through the clearances necessary

A-fluid flowing through the clearance between the tube and the hole in the baffle.

C-fluid flowing through the clearance between the outer tubes and the shells E-fluid flowing through the clearance between the baffle and the shell.

F-fluid flowing through the gap between the tubes because of any pass-partition plates. This is especially significant with a vertical gap.

Because stream A does not bypass the tubes, it is the pressure drop rather than the heater transfer which is affected. Streams C. E and F bypass the tubes, thus reducing the effective



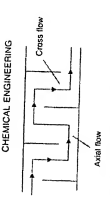


Figure 9.79, Idealised main strenn flow

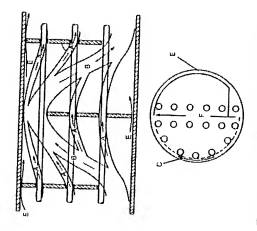


Figure 9.80. Shell-side leakage and by-pass paths [116]

heat transfer area. Stream C, the main bypass stream, is most significant in pull-throughundle units where there is of necessity a large clearance between the bundle and an shell, although this can be reduced by using horizontal scaling strips. In a similar wat the flow of stream F may be reduced by fitting dummy tubes. As an exchanger become fouled, clearances tend to plug and this increases the pressure drop. The whole question of shell-side pressure drop estimation in relation to design procedures is now discussed.

9.9.6. Heat exchanger design

Process conditions

A first-stage consideration in the design process is the allocation of fluids to either shall or tubes and, by and large, the more corrosive fluid is passed through the tubes to reduce the costs of expensive alloys and clad components. Similarly, the fluid with the greater fouling tendency is also usually passed through the tubes where cleaning is easier. Further more, velocities through the tubes are generally higher and more readily controllable at can be adjusted to reduce fouling. Where special alloys are in contact with hot fluids the

And should be passed through the tubes to reduce costs. In addition the shell temperature lowered, thus reducing lagging costs. Passing hazardous materials through the tubes lowered, thus reducing lagging costs. Passing hazardous materials through the tubes seads to greater safety and, because high pressure tubes are cheaper than a high pressure are best handled on the tube-side. In a similar way, where a very low pressure drop is required as in vacuum operation for example, the fluids where a very low pressure drop. Provided the flow is turbulent, a higher heat transfer obtained for a given pressure drop. Provided the flow is turbulent, a higher heat transfer coefficient is usually obtained with a more viscous liquid in the shell because of the more complex flow patterns although, because the tube-side coefficient can be predicted with greater accuracy, it is better to place the fluid in the tubes if turbulent flow in the shell is not possible. Normally, the most economical design is achieved with the fluid with the lower flower flowarte in the shell.

Inselecting a design velocity, it should be recognised that at high velocities high rates of heat transfer are achieved and fouling is reduced, but pressure drops are higher. Normally, the velocity must not be so high as to cause crosion which can be reduced at the tube intention process liquids, velocities are usually 0.3–1.0 m/s in the shell and 1.0–2.0 m/s in the tubes, with a maximum value of 4.0 m/s when fouling must be reduced. Typical water velocities are 1.5–2.5 m/s. For vapours, velocities lie in the range 5–10 m/s with high pressure fluids and 50–70 m/s with vacuum operation, the lower values being used for materials of high molecular weight.

In general, the higher of the temperature differences between the outlet temperature of one stream and the inlet temperature of the other should be 20 deg K and the lower temperature difference should be 5–7 deg K for water coolers and 3–5 deg K when using refrigerated brines, although optimum values can only be determined by an economic analysis of alternative designs.

Similar considerations apply to the selection of pressure drops where there is freedom of choice, although a full economic analysis is justified only in the case of very expensive antis. For liquids, typical values in optimised units are 35 kN/m² where the viscosity is less than 1 mN s/m² and 50–70 kN/m² where the viscosity is 1–10 mN s/m²; for gases, 0.4-0.8 kN/m² for high vacuum operation, 50 per cent of the system pressure at 100–200 kN/m²; and 10 per cent of the system pressure above 1000 kN/m². Whatever pressure drop is used, it is important that erosion and flow-induced tube vibration caused by high velocity, fluids are avoided.

Design methods

His shown in Section 9.9.5 that, with the existence of various bypass and leakage streams in Practical heat exchangers, the flow patterns of the shell-side fluid, as shown in Figure 9.79, are complex in the extreme and far removed from the idealised cross-flow situation discussed in Section 9.4.4. One simple way of using the equations for cross-flow presented in Section 9.4.4, however, is to multiply the shell-side coefficient obtained from these equations by the factor 0.6 in order to obtain at least an estimate of the shell-side coefficient in a practical siluation. The pioneering work of KERN⁽²⁸⁾ and DONOHUE⁽¹¹¹⁾, who used correlations based on the total stream flow and empirical methods to allow for the performance of real exchangers compared with that for cross-flow over ideal tube banks, went much further and,

528

although their early design method does not involve the calculation of bypass and leakage streams, it is simple to use and quite adequate for preliminary design calculations.

The method, which is based on experimental work with a great number of commercial exchangers with standard tolerances, gives a reasonably accurate prediction of heat transfer coefficients for standard designs, although predicted data on pressure drop is less satisfactory as it is more affected by leakage and bypassing. Using a similar approach that for tube-side flow, shell-side heat transfer and friction factors are correlated using a hypothetical shell diameter and shell-side velocity where, because the cross-sectional area for flow varies across the shell diameter, linear and mass velocities are based on the maximum area for cross-flow; that is at the shell equator. The shell equivalent diameter is obtained from the flow area between the tubes taken parallel to the tubes, and the wetted perimeter, as outlined in Section 9.9.4 and illustrated in Figure 9.28. The shells side factors, j_b and j_f , for various baffle cuts and tube arrangements based on the dangiven by KERR⁽³⁸⁾ and LUDWIG⁽¹⁰¹⁾ are shown in Figures 9.81 and 9.82.

The general approach is to calculate the area for cross-flow for a hypothetical row tubes at the shell equator from the equation given in Section 9.4.4:

$$A_s = d_s I_{\theta} C / Y \tag{9.226}$$

where d_s is the shell diameter. I_B is the baffle length and (C'/Y) is the rutio of the clearance between the tubes and the distance between tube centres. The mass flow divided by the area A_s gives the mass velocity G_s , and the linear velocity on the shell-side u_s is obtained by dividing the mass velocity by the mean density of the fluid. Again using the equations in Section 9.4.4, the shell-side equivalent or hydraulic diameter is given by:

For square pitch:
$$d_r = 4(Y^2 - \pi d_0^2)4/\pi d_0 = 1.27(Y^2 - 0.785d_0^2)/d_0^9.227$$

and for triangular pitch : $d_v = 4\{(0.87Y \times Y/2) - (0.5\pi d_o^2/4)/(\pi d_o/2)\}$ = 1.10(Y² - 0.917 d_o^2)/ d_o

(9.228)

Using this equivalent diameter, the shell-side Reynolds number is then:
$$Re_s = G_s' d_s / \mu = u_s d_s \rho / \mu$$
 (9.229)

where G'_j is the mass flowrate per unit area. Hence j_k may be obtained from Figure 9.81. The shell-side heat transfer coefficient is then obtained from a re-arrangement of equation 9.220;

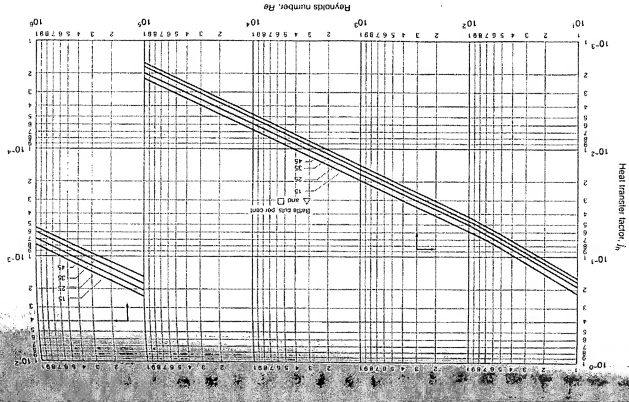
$$Nu = (h_i d_e / k_f) = j_h ReP r^{0.33} (\mu / \mu_s)^{0.14}$$
 (9.230)

In a similar way, the factor j_f is obtained from Figure 9.82 and the pressure drop estimated from a modified form of equation 9.224:

$$-\Delta P_s = 4j_I(d_s/d_e)(I/I_B)(\rho n_s^2)(\mu/\mu_s)^{-0.14}$$
 (9.231)

where $(1/I_B)$ is the number of times the flow crosses the tube bundle = (n+1).

The pressure drop over the shell nozzles should be added to this value although this is usually only significant with gases. In general, the nozzle pressure loss is 1.5 velocity heads for the inlet and 0.5 velocity heads for the outlet, based on the nozzle area or the



rec area between the tubes in the row adjacent to the nozzle, whichever is the least. Kent's method is now illustrated in the following example. may the state of the said of

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Example 9.28
Example 9.28
Graff v. method, design a shell and tube heat exchanger to cool 30 kg/s of buyl alcohol from 370 to 0.008 Kern's method, design a shell and tube heat exchanger at 300 K and leave at 315 K.

Solution

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 $\Pi\Pi$

Since the corrisive, the water will be passed through the tubes.

An a mean temperature of 0.5(370 ± 315) = 343 K, from Table 3. Appendix A1, the thermal capacity of hard stable = 2.90 kJ/kg K and hence:

Heat load = $(30 \times 2.90)(370 - 315) = 4785 \text{ kW}$

Flow of cooling water = 4785/(4.18(315 - 3001) = 76.3 kg/s High leaft capacity of water is 4.18 kHkg K, then:

The logarithmic mean temperature difference,

10,

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$$\theta_n = \{(370 - 315) - (315 - 300)\}/(11(370 - 315)/(315 - 300)\} = 30.7 \text{ deg K}$$

With one shell-side pass and two tube-side passes, then from equation 9.213:

X = (370 - 315)/(315 - 300) = 3.67 and Y = (315 - 300)/(370 - 300) = 0.21

and from Figure 9.75:

Reynolds numbor, Re Shell side (riction factions with segmental balffins (33)

97 32

Baffle cuts, per cent

$$F = 0.85$$
 and $F\theta_m = (0.85 \times 30.7) = 26.1$ deg K

From Table 9.17, an estimated value of the overall coefficient is $U=500~\mathrm{W/m^2 K}$ and hence, the provisional equation 9.212, is:

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$$A = (4785 \times 10^3)/(26.1 \times 500) = 367 \text{ m}^2$$

It is convenient to use 20 mm OD, 16 mm ID tubes, 4.88 m long which, allowing for the tube-sheets, would provide an effective tube length of 4.83 m. Thus:

Surface area of one (ube = $\pi(20/1000) = 0.303 \text{ m}^2$

Number of tubes required = (367/0.303) = 1210

AN TO STATE SIDE (1.25 triangular pitch may be used and, from equation 9.211;

 $1210 = 0.249 (d_b/20)^{2.207}$

$$d_b = 937 \text{ mm}$$

from which

10_S

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168782

Unig a spiliting floating head unit, then, from Figure 9.71, the diametrical clearance between the shell and the tibes = 68 mm and:

Shell diumeter, $d_{\rm v} = (937 \pm 68) = 1005 \text{ mm}$

which approximates to the nearest standard pipe size of 1016 mm.

coefficient Tube-side

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Z £

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e 1 -01

Friction factor, j_f

Z-01

Toward side coefficient may now be calculated using equation 9,218, although here, use will be made of the

Cross-sectional area of one tube = $(\pi/40) \times 16^2 = 201$ num²

Number of tubes/pass = (1210/2) = 605

Thus:

Tube-side flow area = $(605 \times 201 \times 10^{-9}) = 0.122 \text{ m}^2$

Mass velocity of the water = $(76.3/0.122) = 625 \text{ kg/m}^2\text{s}$

Thus, for a mean water density of 995 kg/m3;

Water velocity,
$$u = (625/995) = 0.63$$
 m/s

At a mean water temperature of 0.5(315 \pm 300) = 308 K, viscosity, μ = 0.8 mN s/m² and thermal conductivity, k = 0.59 W/m K.

$$Re = du\rho/\mu = (16 \times 10^{-3} \times 0.63 \times 995)/(0.8 \times 10^{-3}) = 12540$$

$$Pr = C_p \mu / k = (4.18 \times 10^3 \times 0.8 \times 10^{-3})/0.59 = 5.67$$

 $1/d_i = 4.83/(16 \times 10^{-3}) = 302$

Thus, from Figure 9.77, $j_b = 3.7 \times 10^{-3}$, and, in equation 9.218, neglecting the viscosity term:

$$(h_t \times 16 \times 10^{-3})/0.59 = (3.7 \times 10^{-3} \times 12540 \times 5.67^{0.33})$$

h, = 3030 W/m2K

and;

Shell-side coefficient

The baffle spacing will be taken as 20 per cent of the shell diameter or $(1005 \times 20/100) = 201$ mm. The tube pitch = $(1.25 \times 20) = 25$ mm and, from equation 9.226:

Cross-flow area,
$$A_s = \{(25 - 20)/25 | (1005 \times 201 \times 10^{-6}) = 0.040 \text{ m}^2 \}$$

Mass velocity in the shell, $G_t = (30/0.040) = 750 \text{ kg/m}^3$ s Thus:

From equation 9.228;

Equivalent diameter,
$$d_r = 1.10[25^2 - (0.917 \times 20^2)]/20 = 14.2 \text{ mm}$$

At a mean shell-side temperature of 0.5(370 + 315) = 343 K, from Appendix A1;

density of buryl alcohol, $\rho=780 \text{ kg/m}^3$, viscosity, $\mu=0.75 \text{ mN s/m}^2$, hent capacity, $C_p=3.1 \text{ kJ/kg/K}$

and thermal conductivity, k = 0.16 W/m K. Thus, from equation 9.229;

$$Re = G_{\rm t} d_{\rm e}/\mu = (750 \times 14.2 \times 10^{-3})/(0.75 \times 10^{-3}) = 14200$$

 $Pr = C_p \mu / k = (3.1 \times 10^3 \times 0.75 \times 10^{-3})/0.16 = 14.5$

Thus, with a 25 per cent segmental cut, from Figure 9.81: $f_h = 5.0 \times 10^{-3}$ Neglecting the viscosity correction term in equation 9.230:

 $(h_r \times 14.2 \times 10^{-3})/0.16 = 5.0 \times 10^{-3} \times 14200 \times 14.5^{0.31}$

 $h_{\rm c} = 1933 \text{ W/m}^2 \text{K}$

The mean butanol temperature = 343 K, the mean water temperature = 308 K and hence the mean wall temper ature may be taken as 0.5(343 + 308) = 326 K at which $\mu_x = 1.1 \text{ mN s/m}^2$

$$(\mu/\mu_1)^{0.14} = (0.75/1.1)^{0.14} = 0.95$$

showing that the correction for a low viscosity fluid is negligible.

Overall coefficient

The thermal conductivity of cupro-nickel alloys = 50 W/m K and, from Table 9.16, scale resistances will be taken as 0.00020 m²K/W for the water and 0.00018 m²K/W for the organic.

Based on the outside area, the overall coefficient is given by:

HEAT TRANSFER

$$1/U = 1/h_0 + R_0 + x_w/k_w + R_1/(d_0/d_0) + (1/h_0)(d_0/d_0)$$

$$\mathbb{R} = (1/1933) + 0.00020 + [0.5(20 - 16) \times 10^{-3}/50] + (0.00015 \times 20)/16 + 20/(3030 \times 16)$$

$$= 0.00052 \pm 0.00020 \pm 0.00004 \pm 0.000225 \pm 0.000041 = 0.001140 \text{ m}^2 \text{K/W}$$

*and: U = 717 W/m²K

when is well in excess of the assumed value of 500 W/m²K.

pressure drop

On the inbe-side, Re=12450 and from Figure 9.78, $j_f=4.5\times10^{-3}$ Veglecting the viscosity correction term, equation 9.225 becomes:

$$\Delta P_1 = 2(4 \times 4.5 \times 10^{-3} (4830/16) + 1.25)(995 \times 0.63^2) = 5279 \text{ N/m}^2 \text{ or } 5.28 \text{ k/N/m}^2$$

which is low, permitting a possible increase in the number of tube passes. On the shell-side, the linear velocity, $(G_s/\rho) = (750/780) = 0.96$ m/s

Neglecting the viscosity correction term, in equation 9.231: Figure 9.82, when Re = 14200, J_f = 4.6 × 10"

$$-\Delta P_s = (4 \times 4.6 \times 10^{-2})(1005/14.2)(4830/201)(780 \times 0.96^2)$$

= 224950 N/m² or 225 kN/m²

This value is very high and thought should be given to increasing the baffle spacing. If this is doubled, this will reduce the pressure drop by approximately $(1/2)^2 = 1/4$ and:

Since h, & Re 0.3 Cx 116.5.

$$h_o = 1933(1/2)^{0.8} = 1110 \text{ W/m}^2\text{K}$$

which gives an overall coefficient of 561 W/m²K which is still in excess of the assumed value of 500 W/m²K. Further detailed discussion of Kern's method together with a worked example is presented in Volume 6.

Whilst Kern's method provides a simple approach and one which is quite adequate for preliminary design calculations, much more reliable predictions may be achieved by taking into account the contribution to heat transfer and pressure drop made by the various idealised flow streams shown in Figure 9.80. Such an approach was originally taken by TINKER(116) and many of the methods subsequently developed have been based on his model which unfortunately is difficult to follow and tedious to use. The approach has been simplified by Devore $^{(11\tilde{t})}$, however, who, in using standard tolerances for commercial exchangers and a limited number of baffle designs, gives nomographs which enable the method to be used with simple calculators. Devore's method has been further simplified by WEFLER⁽¹¹⁸⁾ who gives an illustrative example. Palen and TABOREK $^{(19)}$ and GranT $^{(120)}$ have described how both Heat Transfer Inc. and Heat Transfer and Fluid Flow Services bave used Tinker's method to develop proprietary computer-based methods.

FUSING Tinker's approach, BELL^(121,122) has described a semi-analytical method, based on work at the University of Delaware, which allows for the effects of major bypass and leakage Steams, and which is suitable for use with calculators. In this procedure, the heat transfer coefficient and the pressure drop are obtained from correlations for flow over ideal tube banks, applying correction factors to allow for the effects of leakage, bypassing and flow

> 4 Ŧ

in the window zone. This approach gives more accurate predictions than Kern's method and can be used to determine the effects of constructional tolerances and the use of sealing tapes. This method is discussed in some detail in Volume 6, where an illustrative example

Sciences Data Unit, ESDU(124), and it gives a useful calculation technique for providing A more recent approach is that offered by WILLS and JOHNSTON (123) who have devel oped a simplified version of Tinker's method. This has been adopted by the Engineering an iterative procedure, and the flowrate of each stream to be obtained. The estimation of realistic checks on 'black box' computer predictions. The basis of this approach is shown in Figure 9.83 which shows fluid flowing from A to B in two streams --- over the tubes in cross-flow, and bypassing the tube bundle -- which then combine to form a single stream. In addition, leakage occurs between the tubes and the baffle and between the baffle and the shell, as shown. For each of these streams, a coefficient is defined which permits the pressure drop for each stream to be expressed in terms of the square of the This method is of especial value in investigating the effect of various shell-to-baffle and mass velocity for that stream. A knowledge of the total mass velocity and the sum of the pressure drops in each zone enables the coefficients for each stream to be estimated by the heat transfer coefficient and the pressure drop is then a relatively simple operation. baffle-to-tube tolerances on the performance of a heat exchanger, both in terms of hear ransfer rates and the pressure losses incurred.

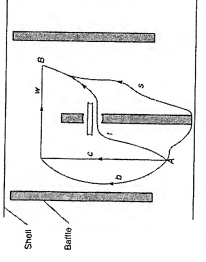


Figure 9.83. Flow streams in the Wills and Johnston method^(1,23)

9.9.7. Heat exchanger performance

One of the most useful methods of evaluating the performance of an existing hear exchanger or to assess a proposed design is to determine its effectiveness n, which is defined as the ratio of the actual rate of heat transfer Q to the maximum rate Qmax that is thermodynamically possible or:

$$I = \frac{Q}{Q_{\text{max}}} \tag{9.232}$$

HEAT TRANSFER

One is the heat transfer rate which would be achieved if it were possible to bring the outlet temperature of the stream with the lower heat capacity to the inlet temperature of the other stream. Using the nomenclature in Figure 9.84, and taking stream I as having the lower value of GCp. then:

$$Q_{\text{max}} = G_1 C_{\mu_1} (T_{11} - T_{21}) \tag{9.233}$$

An overall heat balance gives:

$$Q = G_1 C_{p_1} (T_{11} - T_{12}) = G_2 C_{p_2} (T_{22} - T_{21})$$

Thus, based on stream 1:

$$\eta = \frac{G_1 C_{p_1} (T_{11} - T_{12})}{G_1 C_{p_1} (T_{11} - T_{21})} = \frac{T_{11} - T_{12}}{T_{11} - T_{21}}$$
(9.234)

and, based on stream 2;

$$\eta = \frac{G_2 C_{p_1} (T_{22} - T_{21})}{G_1 C_{p_1} (T_{11} - T_{21})}$$
(9.235)

In calculating temperature differences, the positive value should always be taken.

Example 9.29

A dow of 1 kg/s of an organic liquid of heat capacity 2.0 kJAg K is cooled from 350 to 330 K by a stream of water flowing countercurrently through a double-pipe heat exchanger. Estimate the effectiveness of the unit if the water enters the exchanger at 290 K and leaves at 320 K.

Satution

Heat load,
$$Q = 1 \times 2.0(350 - 330) = 40 \text{ kW}$$

Flow of water,
$$G_{\text{cosl}} = \frac{40}{4.187(320 - 290)} = 0.318 \text{ kg/s}$$

For organic:
$$(GC_p)_{hot} = (1 \times 2.0) = 2.0 \text{ kW/K} (= G_2C_{p_2})$$

$$(GC_p)_{\text{hols}} = (0.318 \times 4.187) = 1.33 \text{ kW/K} (= GC_p)_{\text{min}} = (G_1C_{p_1})$$

equation 9,235:

effectiveness
$$\eta = \frac{2.0(350 - 330)}{1.33(350 - 290)}$$

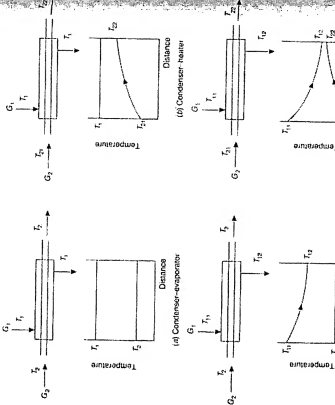
= 0.50

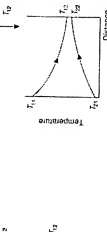
9.9.8. Transfer units

The concept of a transfer unit is useful in the design of heat exchangers and in assessing their performance, since its magnitude is less dependent on the flowrate of the fluids than the treat transfer coefficient which has been used so far. The number of transfer units N is defined by:

$$N = \frac{UA}{(GC_p)_{min}} \tag{9.236}$$

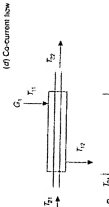
HEAT TRANSFER





Distance

(c) Cooler-evaporator



5

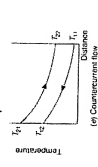


Figure 9.84. Nomenclature for effectiveness of heat exchangers

when its remperature is changed by I deg K. Thus, the number of transfer units gives a measure of the amount of heat which the heat exchanger can transfer. The relation for the where $(G_C_p)_{min}$ is the lower of the two values $G_1C_{p_1}$ and $G_2C_{p_2}$. N is the ratio of the heat in the first of the fluid stream effectiveness of the heat exchanger in terms of the heat capacities of the streams is now Transfer units are also used extensively in the calculation of mass transfer rates in Even for a number of flow conditions. The relevant nomenclature is given in Figure 9.84.

Considering co-current flow as shown in Figure 9.84d, for an elemental area dA of a countercurrent columns and reference should be made to Chapter 10. heat exchanger, the rate of transfer of heat dQ is given by:

$$dQ = U dA(T_1 - T_2) = U dA \theta \tag{9.237}$$

Where \vec{E}_1 and T_2 are the temperatures of the two streams and θ is the point value of the emperature difference between the streams.

maddition:
$$dQ = G_2C_{p_1}dT_2 = -G_1C_{p_1}dT_1$$

Thius: $dT_2 = \frac{dQ}{G_2C_{p_2}}$ and $dT_1 = \frac{-dQ}{G_1C_{p_1}}$
and $dT_1 - dT_2 = d(T_1 - T_2) = d\theta = -dQ\left(\frac{1}{G_1C_{p_1}} + \frac{1}{G_2C_{p_2}}\right)$

Substituting from equation 9.237 for dQ:

$$\frac{\mathrm{d}\theta}{\theta} = -U \mathrm{d}A \left[\frac{1}{G_1 C_{P_1}} + \frac{1}{G_2 C_{P_2}} \right] \tag{9.238}$$

 $\ln \frac{\theta_2}{\theta_1} = -UA \left[\frac{1}{G_1 C_{P1}} + \frac{1}{G_2 C_{P2}} \right]$

$$\ln \frac{T_{12} - T_{23}}{T_{11} - T_{21}} = \frac{UA}{G_1 G_{p_1}} \left[1 + \frac{G_1 G_{p_1}}{G_2 C_{p_2}} \right]$$
(9.239)

$$\Pi G_1 C_{p_1} < G_2 C_{p_2}, G_1 C_{p_1} = (GC_p)_{min}$$

From equation 9,236;

$$\frac{T_{12} - T_{22}}{T_{11} - T_{21}} = \exp\left[-N\left(1 + \frac{G_1C_{P_1}}{G_2C_{P_2}}\right)\right]$$
(9.240)

From equations 9.234 and 9.235:

$$T_{11} - T_{12} = \eta(T_{11} - T_{21})$$

 $T_{22} - T_{21} = \eta \frac{G_1 C_{P_1}}{G_2 C_{P_2}} (T_{11} - T_{21})$

Adding:
$$T_{11} - T_{12} + T_{22} - T_{21} = \eta \left(1 + \frac{G_1 C_{p_1}}{G_2 C_{p_2}} \right) (T_{11} - T_{21})$$

$$1 - \frac{T_{12} - T_{22}}{T_{11} - T_{21}} = \eta \left(1 + \frac{G_1 C_{p_1}}{G_2 C_{p_2}} \right)$$

Substituting in equation 9.240;

$$\eta = \frac{1 - \exp\left[-N\left(1 + \frac{G_1C_{p_1}}{G_2C_{p_2}}\right)\right]}{1 + \frac{G_1C_{p_1}}{G_2C_{p_2}}}$$
 (6)

For the particular case where $G_1C_{p_1} = G_2C_{p_2}$:

$$\eta = 0.5[1 - \exp(-2N)]$$

[NIZ-] (NIZ-) [NIZ-)

For a very large exchanger $(N \to \infty)$, $\eta \to 0.5$. A similar procedure may be followed for *countercurrent flow* (Figure 9.84e), although it should be noted that, in this case, $\theta_1 = T_{11} - T_{22}$ and $\theta_2 = T_{12} - T_{21}$.

The corresponding equation for the effectiveness factor η is then:

$$\eta = \frac{1 - \exp\left[-N\left(1 - \frac{G_1C_{P_1}}{G_2C_{P_2}}\right)\right]}{1 - \frac{G_1C_{P_1}}{G_2C_{P_2}}\exp\left[-N\left(1 - \frac{G_1C_{P_1}}{G_2C_{P_2}}\right)\right]}$$
(9.2)

For the case where $G_1C_{p_1}=G_2C_{p_2}$, it is necessary to expand the exponential terms G_1

$$\frac{N}{N+1} = n$$

In this case, for a very large exchanger $(N \to \infty)$, $\eta \to 1$.

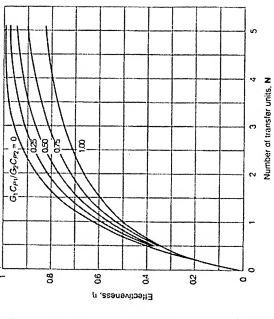
If one component is merely undergoing a phase change at constant temperature, (Figure 9.84b, c) $G_1C_{p_1}$ is effectively zero and both equations 9.241 and 9.243 reduce to

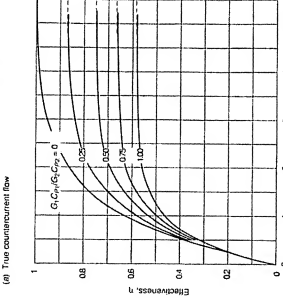
$$\eta = 1 - \exp(-iN)$$

Effectiveness factors η are plotted against number of transfer units N with $(G_1C_{p_1}/G_2C_{p_1})$ as parameter for a number of different configurations by KAYS and LONDON⁽¹²⁵⁾. Examples for countercurrent flow (based on equation 9.235) and an exchanger with one shell pass and two tube passes are plotted in Figures 9.85 σ and σ respectively.

Example 9.30

A process requires a flow of 4 kg/s of puritied water at 340 K to be heated from 320 K by 8 kg/s of unireated water which can be available at 380, 370, 360 or 350 K. Estimate the heat transfer surfaces of one shell pass two tube pass heat exchangers suitable for these duties. In all cases, the mean heat capacity of the water streams is 4.18 kJ/kg K and the overall coefficient of heat transfer is 1.5 kW/m² K.





Number of transfer units, N (b) One shell pass, two-tube pass exchanger

Figure 9.85. Effectiveness of heat exchangers as a function of number of transfer units (125)

Solution

 $GC_p = (8.0 \times 4.18) = 33.44 \text{ kW/K}$ For the untreated water:

 $GC_p = (4.0 \times 4.18) = 16.72 \text{ kW/K}$ For the purified water.

Thus:

 $(GC_p)_{\text{trip}} = 16.72 \text{ kW/K} = G_1C_{p_1}$

 $\frac{G_1G_{p_1}}{G_2G_{p_2}} = \frac{16.72}{33.44} = 0.5$

From equation 9,235;

and:

$$\eta = \frac{4.0 \times 4.18(340 - 320)}{4.0 \times 4.18(T_{11} - 320)}$$

 $(T_{11} - 320)$

Thus η may be calculated from this equation using values of $T_{11}=380,370,360$ or 350 K and then N obtains from Figure 9.85b. The area required is then calculated from:

(equation 9.2 $A = \frac{N(GC_P)_{\text{fitted}}}{U}$

to give the following results:

A (m²)	5.0	9.9	001	200
(-) N	0.45	0.6	6.0	
(<u> </u>	0.33	0.4	0.5	0.67
T ₁₁ (K)	380	370	360	350

Obviously, the use of a higher untreated water temperature is attractive in minimising the area required although in practice any advantages would be offset by increased water costs, and an optimisation procedur would be necessary in obtaining the most effective design.

9.10. OTHER FORMS OF EQUIPMENT

9.10.1. Finned-tube units

Film coefficients

surface on the side with the limiting coefficient. This may be done conveniently by using When viscous liquids are heated in a concentric tube or standard tubular exchanger by This condition also arises with air or gas heaters where the coefficient on the gas side side. It is often possible to obtain a much better performance by increasing the area of will be very low compared with that for the liquid or condensing vapour on the other condensing steam or hot liquid of low viscosity, the film coefficient for the viscous liquid is much smaller than that on the hot side and it therefore controls the rate of heat transfer a finned tube as in Figure 9.86 which shows one typical form of such equipment which may have either longitudinal or transverse fins.

The calculation of the film coefficients on the fin side is complex because each unit. of surface on the fin is less effective than a unit of surface on the tube wall. This arrest occause there will be a temperature gradient along the fin so that the temperature difference



Ingure 9.866. Tube with tadial fins

ficients it is convenient to consider firstly the extended surface as shown in Figure 9. A cylindrical rod of length L and cross-sectional area A and perimeter b is heated a_0 end by a surface at temperature T_1 and cooled throughout its length by a mediun between the fin surface and the fluid will vary along the fin. To calculate the films temperature T_G so that the cold end is at a temperature T_2 .

A heat balance over a length dv at distance x from the hot end gives:

heat in = heat out along rod + heat lost to surroundings

$$-kA\frac{dT}{dx} = \left[-kA\frac{dT}{dx} + \frac{d}{dx} \left(-kA\frac{dT}{dx} \right) dx \right] + hb\,dx(T - T_G)$$

Or:

where h is the film coefficient from fin to surroundings.

Writing the temperature difference $T-T_G$ equal to θ :

$$kA\frac{d^2T}{dr^2}dx = hb\,dx\,\theta$$

Since T_G is constant, $d^2T/dx^2 = d^2\theta/dx^2$.

$$\frac{\mathrm{d}^2\theta}{\mathrm{d} r^2} = \frac{hb}{kA}\theta = m^2\theta. \quad \left(\text{where } m^2 = \frac{hb}{kA}\right)$$

Thus:

and:

 $\theta = C_1 e^{mx} + C_2 e^{-mx}$

In solving this equation, three important cases may be considered:

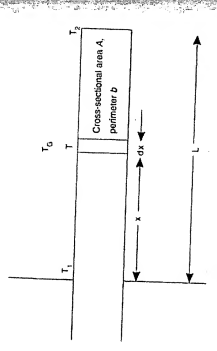


Figure 9.87. Heat flow in rod with heat loss to surroundings

(a) A long rod with temperature falling to that of surroundings, that is $\theta = 0$ when $x = \infty$

HEAT TRANSFER

b) A short and from which heat loss from its end is neglected.

At the hot end:
$$x = 0$$
, $\theta = \theta_1 = C_1 + C_2$

At the cold end:
$$x = L$$
, $\frac{d\theta}{dr} = 0$

$$0 = C_1 m e^{mt} - C_2 m e^{-mt}$$

$$\theta_1 = C_1 + C_1 e^{2mL}$$

$$\theta_1 = C_1 + C_2 e^{2mL}$$

$$C_1 = \frac{\theta_1}{1 + e^{2mL}}, \quad C_2 = \frac{\theta_1}{1 + e^{-2mL}}$$
$$\theta = \frac{\theta_1 e^{mt}}{1 + e^{2mL}} + \frac{\theta_1 e^{2mL} e^{-mt}}{1 + e^{2mL}}$$

$$= \frac{\theta_1}{1 + e^{2mL}} [e^{m_1} + e^{2mL}e^{-m_1}]$$
 (9.249)

$$\frac{\theta}{\theta_1} = \frac{e^{-mL}e^{mx} + e^{mL}e^{-mx}}{e^{-mL} + e^{mL}}$$

$$\frac{\theta}{\theta_1} = \frac{\cosh m(L - x)}{\cosh mL}$$

This may be written:

(9.250)

c) More accurately, allowing for heat loss from the end:

At the hot end:
$$x = 0$$
, $\theta = \theta_1 = C_1 + C_2$

At the cold end:
$$x = L$$
, $Q = hA\theta_{n=L} = -kA\left(\frac{d\theta}{dx}\right)_{x=L}$

The defermination of C₁ and C₂ in equation 9.247 then gives:

$$\theta = \frac{\theta_1}{1 + Je^{-2mL}} (Je^{-2mL}e^{mx} + e^{-mx})$$
 (9.251)

or again, noting that
$$\cosh x = \frac{1}{2}(e^x + e^{-x})$$
 and $\sinh x = \frac{1}{2}(e^x - e^{-x})$:

$$\frac{\theta}{\theta_1} = \frac{\cosh m(L - x) + (h/mk) \sinh m(L - x)}{\cosh mL + (h/mk) \sinh mL}$$
(9.252)

The hear loss from a finned tube is obtained initially by determining the heat flow into the base of the fin from the tube surface. Thus the heat flow to the root of the fin is: $Q_f = -kA \left(\frac{\mathrm{d}T}{\mathrm{d}x} \right)_{x=0} = -kA \left(\frac{\mathrm{d}\theta}{\mathrm{d}x} \right)_x$

For case (a):

$$Q_f = -kA(-m\theta_1) = kA\sqrt{\frac{hb}{kA}\theta_1} = \sqrt{hbkA}\theta_1$$

For case (b):

$$Q_f = -kAm\theta_1 \frac{1 - e^{2mt}}{1 + e^{2mt}} = \sqrt{hbkA} \, \theta_1 \tanh mL$$

For case (c):

$$Q_f = \sqrt{hbkA}\,\theta_1 \left(\frac{1 - Je^{-2mL}}{1 + Je^{-2mL}}\right)$$

These expressions are valid provided that the cross-section for heat flow remain constant. When it is not constant, as with a radial or tapered fin, for example, the temperature distribution is in the form of a Bessel function (120).

If the fin were such that there was no drop in temperature along its length, then the maximum rate of heat loss from the fin would be:

$$Q_{f \, \text{max}} = bLh\theta_1$$

The fin effectiveness is then given by $Q_f/Q_{f \, \text{max}}$ and, for case (b), this becomes,

$$\frac{kAm\theta_1\tanh mL}{bLh\theta_1} = \frac{\tanh ml}{mL}$$

Example 9.31

In order to measure the temperature of a gas flowing through a copper pipe, a thermometer pocket is fine perpendicularly through the pipe wall, the open end making very good contact with the pipe wall. The pocket is made of copper tube, 10 mm o.d. and 0.9 mm wall, and it projects 75 mm into the pipe. A thermocouple welded to the bottom of the tube and this gives a reading of 475 K when the wall temperature is at 365 K life, coefficient of heat transfer between the gas and the copper tube is 140 W/m² K, calculate the gas remarked the transfer between the gas and the copper tube is 140 W/m² K, calculate the gas remperature. The thermal conductivity of copper may be taken as 350 W/m K. This arrangement is shown in Figure 9.88.

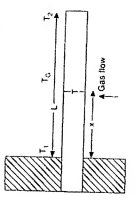


Figure 9.88. Heat transfer to thermometer pocket

Solution

If θ is the temperature difference $(T-T_G)$, then:

$$\theta = \theta_1 \frac{\cosh m(L - x)}{\cosh mL}$$

$$\theta = \frac{\theta_1}{\cosh mL}$$

m, = th

$$b = \pi \times 0.010$$
 m. tube i.d. = 8.2 mm or 0.0082 m

shee the perimeter.

nal area of metal:
$$A = \frac{\pi}{4} (10.0^2 - 8.2^3) = 8.19 \,\pi \,\text{mm}^2$$
 or $8.19 \,\pi \times 10^{-6} \,\text{m}^2$
 $\frac{\pi}{4} (140 \times 0.010 \pi)$

$$m^2 = \frac{(140.7 \text{ OM/S})}{(350 \times 8.19\pi \times 10^{-6})} = 488 \text{ m}^{-2}$$

 $m = 22.1 \text{ m}^{-1}$

$$\theta_1 = T_G - 365$$
, $\theta_2 = T_G - 475$

$$\frac{\theta_1}{\theta_2} = \cosh mL$$

$$\frac{T_G - 365}{T_G - 475} = \cosh(22.1 \times 0.075) = 2.72$$

$$T_G = 530 \text{ K}$$

ple 9.32

and the fitted with transverse circular steel fins of constant cross-section has the following specification:

tube e.d.:
$$d_2 = 54.0 \text{ mm}$$
 in diameter $d_1 = 70.0 \text{ mm}$

fin thickness:
$$w = 2.0$$
 mm number of fins/metre run = 230

*Determine the heat loss per metre run of the tube when the surface temperature is 370 K and the temperature of the surroundings 280 K. The heat transfer coefficient between gas and fin is 30 W/m² K and the thermal conductivity of steel is 43 W/m K.

- Solving

Assuming that the height of the fin is small compared with its circumference and that it may be treated as a traight fin of length $(\pi/2)(d_1+d_2)$, then:

 $A = \frac{\pi(d_1 + d_2)w}{r}$, i.e. the average area at right-angles to the heat flow

$$m = \left(\frac{hb}{kA}\right)^{0.5} = \left\{\frac{h\pi(d_1 + d_2)}{[k\pi(d_1 + d_2)w/2]}\right\}^{0.5}$$
$$= \left(\frac{2h}{kw}\right)^{0.5}$$
$$= \left(\frac{2}{kw}\right)^{0.5}$$
$$= \left(\frac{2 \times 30}{43 \times 0.002}\right)^{0.5}$$

From equation 9.254, the heat flow is given for case (b) as:

$$Q_f = mkA\theta_1 \frac{e^{2mt} - 1}{1 + e^{2mt}}$$

In this equation;

$$A = \frac{(\pi(70.0 + 54.0) \times 2.01}{2} = 390 \text{ mm}^2 \text{ or } 0.00039 \text{ m}^2$$

$$L = \frac{d_1 - d_2}{2} = 8.0 \text{ mm} \text{ or } 0.008 \text{ m}$$

$$mL = 26.42 \times 0.008 = 0.211$$

$$\theta_1 = 370 - 280 = 90 \text{ deg K}$$

$$Q_f = \frac{26.42 \times 43 \times 3.9 \times 10^{-4} \times 90(c^{0.422} - 1)}{2.000}$$

$$= \frac{39.9 \times 0.525}{2.525} = 8.29 \text{ W per tin}$$

The heat loss per metre run of tube = $8.29 \times 230 = 1907 \ \mathrm{W/m}$

In this case, the low value of mL indicates a fin efficiency of almost 1.0, though where mL tends to 110.

Practical data

A neat form of construction has been designed by the Brown Fintube Company of America. On both prongs of a hairpin tube are fitted horizontal fins which films. fin side is balanced by the increase in surface. An indication of the surface obtained is concentric tubes, joined at the base of the hairpin. Units of this form can be convenient. arranged in banks to give large heat transfer surfaces. It is usual for the extended surface to be at least five times greater than the inside surface, so that the low coefficient on the

Table 9.20. Data on surface of finned tube units^(a)

γ γ γ γ γ απ γ γ απ	inpe (m?/m)	Height of fin		0.5 in 1 1 in 1	62 2363 2363		4.97	86 649T	. 1.70e/2002220e/200220
	Surface of finned pipe (m ² /m)		1	.0.4 mm 0.5	0.689 1.3	1.167 2.164			
	(m ²	Height of fin	12.7 mm		0.385	0.660	0.863	1.066	
		Number	of fins		2 9 8	25.52	28.4	9	
	Outside surface	of pipe	(ft2/ft length)	0.263		0.497		***************************************	
	Ours		(m²/m)	0.08	2	21.0		(a) Brown Fintube Company,	
	Pipe size outside dimester		(ii)		-	}		en Fintub	
â	2 2 2			9.0	48.3			(a) Brow	

length of 6.6 m, height of 0.34 m, and width of 0.2 m. The free area for flow on the fin side is 2.645 mm² against 1.320 mm² on the inside; the ratio of the transfer surface or A typical hairpin unit with an effective surface on the fin side of 9.4 m² has an overall

HEAT TRANSFER

The fin side film coefficient ht has been expressed by plotting:

$$\frac{h_f}{C_p G'} \left(\frac{C_p \mu}{k} \right)^{2/3} \left(\frac{\mu}{\mu_s} \right)^{-0.14} \quad \text{against} \quad \frac{d_s G'}{\mu}$$

wherether is based on the total finside surface area (fin and tube), G' is the mass rate of flow per unit area, and d_e is the equivalent diameter, or

de intaliguetted perimeter for flow (fin + outside of tube + inner surface of shell tube)

6.1 milength: the finned tubes were inserted inside tubes 90 mm inside diameter. With seam on the tube side, and tube oils and kerosene on the fin side, the experimental data Experimental work has been carried out with exchangers in which the inside tube was 48 mm outside diameter and was fitted with 24, 28, or 36 fins (12.5 mm by 0.9 mm) in were well correlated by plotting:

$$\frac{h_f}{C_\rho G'} \left(\frac{C_\rho \mu}{k}\right)^{2/3} \left(\frac{\mu}{\mu_s}\right)^{-0.14} \quad \text{against} \quad \frac{d_\rho G'}{\mu}$$

Typical values were:

$$\frac{h_f}{C_p G'} \left(\frac{C_p \mu}{k}\right)^{2/3} \left(\frac{\mu}{\mu_s}\right)^{-0.14} = 0.25 \quad 0.055 \quad 0.012 \quad 0.004$$

$$\frac{d_e G'}{\mu} = 1 \quad 10 \quad 100 \quad 1000$$

Some indication of the performance obtained with transverse finned nubes is given in Table 9.21. The figures show the heat transferred per unit length of pipe when heating air on the fin side with steam or hot water on the tube side, using a temperature difference of 100 deg K. The results are given for three different spacings of the fins.

	Tuble 9	Table 9.21. Data on finned tubes	finned tubes		
Inside diam, of tube Outside diam, of fin	19 mm 64 mm	25 mm 70 mm	38 mm 100 mm	50 mm	75 mm 140 mm
No. of fins/m run	Heal	Heat transferred (kW/m)	W/m)		
80 80 100	0.47 0.49 0.54	0.63 0.64 0.69	0.37 1.02 1.14	1.07	E 4 8
Inside diam, of tube Outside diam, of fin	a in 2½ in	Lin	3. 5. in in	2 in 4 th in	3 in S ¹ in
No. of fins/ft run	Heat	Hear transferred (Bru/h ft)	uh ft)		
87.8	485 505 565	650 665 720	, 1060 1960 190	1115 1170 1295	1440 1495 1655

(Data taken from catalogue of G. A. Harvey and Co. Ltd. of London.)

A series of plate type heat exchangers which present some special features was developed by the APV Company. The general construction is shown in Figure 9.89 where shows an Alfa-Laval exchanger and from which it is seen that the equipment consists a series of parallel plates held firmly together between substantial head frames. The plate one-piece pressings, frequently of stainless steel, and are spaced by rubber sea guskets cemented into a channel around the edge of each plate. Each plate has a num of troughs pressed out at right angles to the direction of flow and arranged so that interlink with each other to form a channel of constantly changing direction and seen With normal construction the gap-between the plates is 1.3–1.5 mm. Each liquid fit in alternate spaces and a large surface can be obtained in a small volume.

Because of the shape of the plates, the developed area of surface is appreciably great than the projected area. This is shown in Table 9.22 for the four common sizes of plates.

	Table 9.22. Plate areas	Plate a	reas	
	Projected area	area	Developed are	ed are
Plate type	, a	ft.	m,	1
НТ	0.09	1.00	0.13	-
HX	0.13	1.45		500
Ξ	0.27	2.88	0.35	*
¥	0.36	3.85	0.43	30
-				

A high degree of turbulence is obtained even at low flowrates and the high heat train coefficients obtained are illustrated by the data in Table 9.23. These refer to the heat of cold water by the equal flow of hot water in an HF type exchanger (aluminium copper), at an average temperature of 310 K.

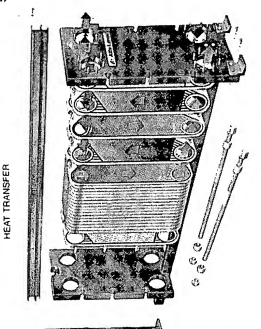
Table 9.23. Performance of plate-type exchanger type HF

Hear tran	ransferred per plate	Water 1	flow .	U based on developed	developed area
W/K	Btu/h *F	lls.	Wieg	kW/m²-K	Bach ft2 'F
1580 2640 2640	3000 4000 5 000	0.700 1.075 1.580	\$50 850 1250	07.£ +0.4 €1.3	650 870 1080

(Courtesy of the APV Company.)

Using a stainless steel plate with a flow of 0.00114 m³/s, the heat transferred is 760 W/K for each plate.

The high transfer coefficient enables these exchangers to be operated with very small temperature differences, so that a high heat recovery is obtained. These units have been particularly successful in the dairy and brewing industries, where the low liquid capacity and the close control of temperature have been valuable features. A further advantage is that they are easily dismantled for inspection of the whole plate. The necessity for the long gasket is an inherent weakness, but the exchangers have been worked successfully



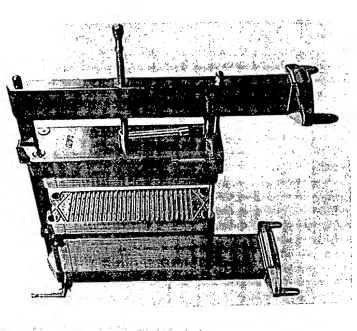


Figure 9.89, (a) Alfa-Laval plate heat exchanger (b) APV plate heat echanger

up to 423 K and at pressures of 930 kN/m². They are now being used in the process and gas industries with solvents, sugar, acetic acid, ammoniacal liquor, and so on,

9.10.3. Spiral heat exchangers

A spiral plate exchanger is illustrated in Figure 9.90 in which two fluids flow three the channels formed between the spiral plates. With this form of construction the ven may be as high as 2.1 m/s and overall transfer coefficients of 2.8 kW/m2 K are frequ obtained. The size can therefore be kept relatively small and the cost becomes compa or even less than that of shell and tube units, particularly when they are fabricated alloy steels.

tube. The construction overcomes problems of differential expansion rates of the tube. pass counterflow heat exchanger with fixed tube plates distinguished by the spiral winds and the shell and the characteristics of the design enable the unit to perform wellser A further design of spiral heat exchanger, described by NEIL(127), is essentially a sin coil such that the inner heat transfer coefficient is 1.92 times greater than for a stra of the tubes, each consisting, typically of a 10 mm o.d. tube wound on to a 38 mm superheated steam where the combination of counterflow, high surface area per volume of shell and the high inside coefficient of heat transfer enables the superi be removed effectively in a unit of reasonable size and cost,

9.10.4. Compact heat exchangers

Advantages of compact units

products and plant problems associated with drilling rigs have, however, all prompted the and fin heat exchangers. The relation between various types of exchanger, insterms presents construction problems and a more realistic approach is to use plate orange from REDMAN(128), In order to obtain a thermal effectiveness in excess of 90 percent countercurrent flow is highly destrable and this is not easily achieved in shell and the development and increased application of compact heat exchangers. Here compacts is a matter of degree, maximising the heat transfer area per unit volume of exchan velocities. Because of the baffle design on the shell-side, the flow at the best may Jescribed as cross-flow, and the situation is only partly redeemed by having a train of the heat transfer area per unit volume of exchanger is shown in Figure 9.91 and as on offshore rigs where space (costing as much as £120,000/m²) is of paramon and this leads to narrow channels. Making shell and tube exchangers more comp able. One important application of compact heat exchangers is the cooling of man units which often have a number of tube-side passes in order to maintain reaso in energy conservation schemes, the general move from bulk chemicals to value exchangers which is an expensive solution. Again in dealing with high value added In general, heat exchanger equipment accounts for some 10 per cent of the cost of a level at which there is no great incentive for innovation. Trends such as the icts. which could well be heat sensitive, a more controllable heat exchanger than and tube unit, in which not all the fluid is heated to the same extent, might be mportance

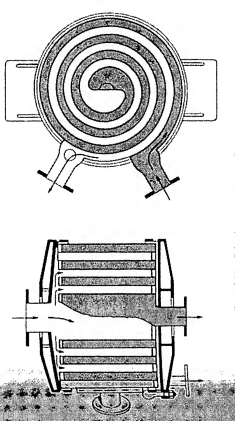


Figure 9.90. Spiral heat exchanger (a) Flow paths

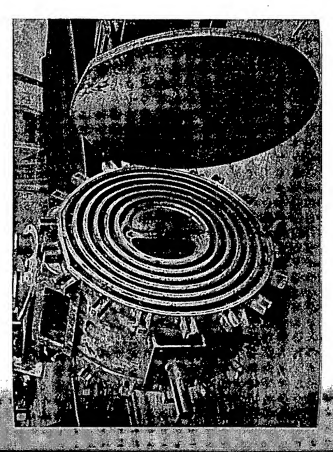


Figure 9.90. Spiral plate exchanger (b) with cover removed

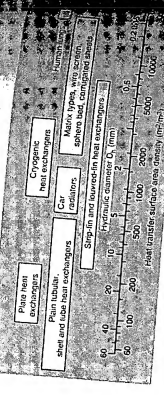


Figure 9.91. Surface area as a function of volume of exchanger for differentives.

Plate and fin exchangers

Plate and his heat exchangers, used in the motor, aircraft and transportindusing years, are finding increased application in the processing industries and in natural gas liquefaction, cryogenic air separation, the production of oldens beparation of hydrogen and carbon monoxide. Potential applications of oldens by GREGORY(129), the concept is that of flat plates of metal, usually alimit plates, and the space between each pair of plates, apportioned to each fluid between the heat transfer. Bars at the edges of the plates retain each fluid between the heat transfer and pressure drop requirements, is known as a layer. The heat designs are shown in Figure 9.92.

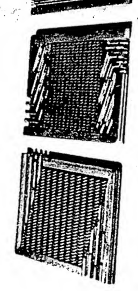


Figure 9.92. Plate and fin exchangers

There are four basic forms of corrugation: plain in which a sheet of metal is come in the simplest way with fins at right angles to the plates; serrated where each fins sideways every 9.5 mm to give a zig-zag path; and perforated corrugation, a changed for a plain corrugation made from perforated material. Each stream to be bear

HEAT TRANSFER

of largers and different corrugation heights, different corrugation types, different of largers and different entry and exit of largers and different entry and exit.

In surface in the hot fluid (or fluids) can vary widely from that in the cold fluid is unrealistic to quote surface areas for either side, as in shell and tube units.

It is unrealistic to quote surface areas for either side, as in shell and tube units.

on the general approach is to obtain the term (HA) for each stream, sum these stream, and determine an overall value of (HA) given by:

$$\frac{1}{(hA)_{nv}} = \frac{1}{(hA)_{v}} + \frac{1}{(hA)_{v}}$$
(9.257)

and a refer, respectively, to overall, hot-side and cold-side values,

ed-circuit exchangers

Configured devices such as small diameter tubes, fins, tube inserts and porous boiling surfaces and tube heart ensemble of improve the surface density and heat transfer coefficients in shell and tube heart extrangers and yet these are not universally applicable and, in general, such units repain essentially bulky. Plate and fin exchangers have either limited fluid compatibility or influed finefficiency, problems which are overcome by using printed-circuit exchangers as essaiped by Joinston and these fluid divergence in the surface destricts of printed circuits. These plates are diffusion-bonded together to form heat exchange cores of the required capacity. Equipheders are attached to the core faces and sometimes the assembly is encapsulated. Passages are (typically 0.3–1.5 mm deep giving surface areas of 1000–5000 m²/m³, an wher of magnitude higher than surface densities in shell and tube designs; and in addition like fine passages tend to sustain relatively high heat transfer coefficients, undiminished in inefficiencies, and so less surface is required.

In designing a unit, each side of the exchanger is independently tailored to the duty function indithe exchanger effectiveness (discussed in Section 9.9.4) can range from 2-5 cent to values in excess of 98 per cent without fundamental design or construction oblines arising. Countercurrent, co-current and cross-flow contacting can be employed dividually or in combination.

A note of caution on the use of photo-etched channels has been offered by tamestaw (31) who points out that the system is attractive in principle provided that

**RANSHAW(31) who points out that the system is attractive in principle provided that we'ver practical problems such as fouling are not encountered. With laminar flow in galnees with a mean plate spacing of 0.3-1 mm, volumetric heat transfer coefficients of "AW/m" K have been obtained with modest pressure drops. Such values compare with 0.2 MW/m" K for shell and tube exchangers and 1.2 MW/m" K for plate heat exchangers.

9.105. Scraped-surface heat exchangers

In eases where a process fluid is likely to crystallise on cooling or the degree of fouling is very ligh or indeed the fluid is of very high viscosity, use is often made of scraped-surface hat exchangers in which a rotating element has spring-loaded scraper blades which wipe

fixed or variable since both rigid and hinged blades may be used. The process liquid continuously spread in a thin layer over the vessel wall and it moves through the device either by the action of gravity or that of the agitator or of both. A tapered or helical arrangement is to connect several sections in series or to install several pipes within common shell. Scraped-surface units of this type are used in paraffin-wax plants and of the inside surface of a tube which may typically be 0.15 m in diameter. Double pipe involved. In such a device the clearance between the agitator and the wall may be enter construction is often employed with a jacket, say 0.20 m in diameter, and one comme to which the thin-film device is especially suited because of the very short residence times evaporating viscous or heat-sensitive materials under high vacuum. This is an application agitator produces longitudinal forces on the liquid.

the rate of cooling depends very much on the effectiveness of the scrapers, and quark overall coefficients of heat transfer ranging from 15 W/m² K with a poorly fitting scrapers to 90 W/m² K where close fitting scrapers remove the wax effectively from the chilled In describing chillers for the production of wax distillates, NELSON(133) points out that

clearance between the agitator and the cynnuca wan survey. Typical applications of the the radial hydrodynamic force of the liquid on the scraper. Typical applications of the scraper. the cylinder wall by the hydrodynamic action of the fluid on the agitator and by centrifug. used extensively in the food processing industries and also in the manufacture of greater and detergents. As the blades are free to move, the clearance between the blades and uswall varies with operating conditions and a typical installation may be 75-100 mm in as agitators and, in general, these are fixed clearance devices used for high viscosing The Votator design has two or more floating scraper-agitators which are forced against action; the blades are loosely attached to a central shaft called the mutator. The volatories the scrupers are held against the wall by leaf springs, and again there is a variable. heat exchangers and chemical reactors employ helical ribbons, augers or twisted tapes. diameter and 0.6-1.2 m long. In the spring-loaded type of scraped-surface heat exchanger the units are 0.15-0.3 m in diameter and up to 12 m long. Some of the more specialised materials. There is no general rule as to maximum or minimum dimensions since cath application is a special case.

where the process fluid changes from a thin liquid to a paste and finally to a powder. This was confirmed by LAUGHLIN(134) who has presented operating data on a system One of the earliest investigations into the effectiveness of scrapers for improving her was negligible, cooling times for more viscous materials could be considerably reduced. HOULTON(135), making tests on the votator, found that back-mixing was negligible may quote overall heat transfer coefficients for each food tested. Using a liquid-full system. SKELLAND et al. (137-139) have proposed the following general design correlation for the transfer was that of HUGGINS(133) who found that although the improvement with water and LINEBERRY(136) who, in addition to discussing the operation and uses of the votator. some useful data on a number of food products have been obtained by BOLANOWSE

$$\frac{hd_v}{k} = c_1 \left(\frac{C_\rho \mu}{k} \right)^{C_2} \left(\frac{(d_v - d_r)u\rho}{\mu} \right) \left(\frac{d_v N}{u} \right)^{0.82} \frac{(d_r)}{d_v} \right)^{0.55} \tag{9.25}$$

 $\frac{1}{2}0.039$ and $c_2 = 0.70$. In this correlation d_v is the diameter of the vessel, d_v is the where for cooling viscous liquids $c_1 = 0.014$ and $c_2 = 0.96$, and for thin mobile liquids danter of the rotor and u is the average axial velocity of the liquid. This correlation only be applied to the range of experimental data upon which it is based, since h will not approach zero as n_B , d_r , $(d_v - d_r)$, N and u approach zero. Reference to the use the wotator for crystallisation is made in Volume 2, Chapter 15.

The majority of work on heat transfer in thin-film systems has been directed towards okuining data on specific systems rather than developing general design methods, although Borner (140-142) have developed the following correlations for heating without change

$$\frac{hd_n}{k} = Nu = 0.018Re^{i0.6}Re^{0.46}p_r^{0.87}\left(\frac{d_v}{l}\right)^{0.48}(n_B)^{0.24} \tag{9.259}$$

, and for evaporation:

$$\frac{hd_v}{k} = Nu = 0.65Re^{-6.43}Re^{0.25}p_r^{0.3}(n_g)^{0.33}$$
 (9.26)

from both of these equations, it will be noted that the heat transfer coefficient is not a function of the temperature difference. Here $Re'' = (d_v^2 N \rho/\mu)$ and $Re = (ud_v \rho/\mu)$, where A is the tube diameter and u is the average velocity of the liquid in the film in the axial direction.

all is also of significance that the agitation suppresses nucleation in a fluid which might otherwise deposit crystals.

9.11. THERMAL INSULATION

9,11.1. Heat losses through lagging

ndiation, conduction, and convection. The loss by radiation is a function of the fourth power of the absolute temperatures of the body and surroundings, and will be small for low A had reaction or storage vessel or a steam pipe will lose heat to the atmosphere by Artista very poor conductor, and the heat loss by conduction will therefore be small except possibly through the supporting structure. On the other hand, since convection turents form very easily, the heat loss from an unlagged surface is considerable. The conservation of hear, and hence usually of total energy, is an economic necessity, and some form of lagging should normally be applied to hot surfaces. Lagging of plant operating a fightemperatures is also necessary in order to achieve acceptable working conditions temperature differences but will increase rapidly as the temperature difference increases. in the vicinity. In furnaces, as has already been seen, the surface temperature is reduced substantially by using a series of insulating bricks which are poor conductors.

is a yery good insulator though it becomes charred at moderate temperatures and is demail conductivity and that it should suppress convection currents. The materials that used mainly in refrigerating plants. Eighty-five per cent magnesia is widely used for seme steam pipes and may be applied either as a hot plastic material or in preformed The two main requirements of a good lagging material are that it should have a low are frequently used are cork, 85 per cent magnesia, glass wool, and vermiculite. Cork retions. The preformed sections are quickly fitted and can frequently be dismantled and re-used whereas the plastic material must be applied to a hot surface and cannot re-used. Thin metal sheeting is often used to protect the lagging.

The rate of heat loss per unit area is given by:

total temperature difference total thermal resistance

For the case of heat loss to the atmosphere from a lagged steam pipe, the there resistance is due to that of the condensate film and dirt on the inside of the pipe has the pipe wall, that of the lagging, and that of the air film outside the lagging The unit length of a lagged pipe:

$$\frac{Q}{l} = \Sigma \Delta T / \left[\frac{1}{h_1 \pi d} + \frac{x_n}{k_n \pi d_n} + \frac{x_l}{k_1 \pi d_m} + \frac{1}{(h_r + h_l)\pi d_s} \right]$$

where d is the inside diameter of pipe, d_w the mean diameter of pipe wall, d_w logarithmic mean diameter of lagging, d_s the outside diameter of lagging, x_s , x_1 are pipe wall and lagging thickness respectively, k., k, the thermal conductivity of the wall and lagging, and h_t , h_r , h_c the inside film, radiation, and convection coefficient

Example 9.33

A steam pipe, 150 mm i.d. and 168 mm o.d., is carrying steam at 444 K and is lagged with 50 mm of cent magnesia. What is the heat loss to air at 294 K?

Solution

In this case;

$$d = 150 \text{ mm}$$
 or 0.150 m
 $d_0 = 168 \text{ mm}$ or 0.168 m
 $d_x = 159 \text{ mm}$ or 0.159 m
 $d_1 = 268 \text{ mm}$ or 0.268 m

The coefficient for condensing steam together with that for any scale will be taken as 8500 Wint K. Co.

 d_m , the log mean of d_o and $d_s = 215$ mm or 0.215 m.

The temperature on the outside of the lagging is estimated at 31.4 K and $(h_r + h_0)$ will be taken 45 W/m K, and k1 as 0.073 W/m K. 10 W/m² K

The thermal resistances are therefore;

$$\frac{s_w}{k_w \pi d_w} = \frac{8500 \times \pi \times 0.150}{0.009} = 0.09025$$

$$\frac{s_w}{k_w \pi d_w} = \frac{0.009}{45 \times \pi \times 0.159} = 0.00040$$

$$\frac{s_0}{k_0 \pi d_m} = \frac{0.039}{0.073 \times \pi \times 0.215} = 1.013$$

$$\frac{1}{(b_w + b_v)\pi d_w} = \frac{1}{10 \times \pi \times 0.268} = 0.119$$

The first two terms may be neglected and hence the total thermal resistance is 1.132 m K/W.

HEAT TRANSFER

 $\frac{1}{100}$ Lead loss per metre length = (444 - 294)/1.132 = 132.5 W/m (from equation 9.261). The temperature on the outside of the lagging may now be checked as follows:

$$\frac{\Delta T(\text{lagging})}{\Sigma \Delta T} = \frac{1.013}{1.132} = 0.895$$

$$\Delta T(\text{lagging}) = 0.895(444 - 294) = 134 \text{ deg K}$$

This the temperature on the cutside of the lagging is (441 – 134) = 310 K, which approximates to the source value.

$$h_t = \frac{[0.9 \times 5.67 \times 10^{-8}(310^2 - 294^4)]}{(310 - 294^3)} = 7.40 \text{ W/m}^2 \text{ K}$$

From Table 9.5 for air (Gr $Pr = 10^4 - 10^9$), $\alpha = 0.25$ and C'' = 1.32. Substituting in equation 9.105 (putting I = diameter = 0.268 m):

$$h_c = C''(\Delta T)^n t^{3n-4} = 1.32 \left[\frac{310 - 294}{0.268} \right]^{0.25} = 3.67 \text{ W/m}^2 \text{ K}$$

Thus $(k_0^2 + k_0^2) = 11.1$ W/m² K, which is close to the assumed value. In practice it is rare for forced throughout currents to be obsent, and the heat loss is probably higher than this value. If the pipe were unlagged. $(h_1 + h_2)$ for $\Delta T = 150$ K would be about 20 W/m² K and the hear loss would

$$\frac{Q}{I} = (h_t + h_c)\pi d_s \Delta T$$

$$= (20 \times \pi \times 0.168 \times 150) = 1584 \text{ W/m}$$
1.58 kW/m

Index these conditions it is seen that the heat loss has been reduced by more than 90 per cent by the addition at stomm thickness of lagging,

8.11.2. Economic thickness of lagging

Mann is suggested for pipes less than 75 mm diameter and 50 mm for pipes up to e an optimum thickness when further increase does not save sufficient heat to justify be too strongly stressed that some lagging everywhere is better than excellent lagging Misome places and none in others. For temperatures of 373-423 K, and for pipes up to Min diameter, LYLE⁽¹⁴³⁾ recommends a 25 mm thickness of 85 per cent magnesia facesing the thickness of the lagging will reduce the loss of heat and thus give a saving the tost in general the smaller the pipe the smaller the thickness used, though it cannot aging and 50 mm for pipes over 230 mm diameter. With temperatures of 470-520 K Table operating costs. The cost of the lagging will increase with thickness and there will

911.3. Critical thickness of lagging

hanceages. Although the outside area from which heat is lost to the surroundings also makes giving rise to the possibility of increased heat loss. It is perhaps easiest to Whe thickness of the lagging is increased, resistance to heat transfer by thermal conduchat of the lagging as acting as a fin of very low thermal conductivity. For a cylindrical

coefficient, and r is the outside diameter of the pipe. In practice, this situation is it. pipe. there is the possibility of heat losses being increased by the application of lagging only if hr/k < 1, where k is the thermal conductivity of the lagging, h is the outside to arise only for pipes of small diameters.

The heat loss from a pipe at a temperature $T_{\mathcal{S}}$ to surroundings at temperature $T_{\mathcal{A}}$ function of x, as shown in Figure 9.93. The rate of heat loss Q from a length l of pur is given by equation 9.262, by considering the heat loss from the outside of the lagging from a constant value T_S at its inner radius r, to an outside temperature T_L which considered. Heat flows through lagging of thickness x across which the temperature fail and by equations 9.263 and 9.264, which give the transfer rate by thermal conduct through the lagging of logarithmic mean radius r_m :

$$Q = 2\pi l(r+x)(T_L - T_A)h$$

$$Q = \frac{2\pi lr_m}{x}k(T_S - T_L)$$

$$= \frac{2\pi lk}{\ln\frac{r+x}{r}}(T_S - T_L)$$

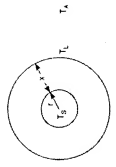


Figure 9.93. Critical lagging thickness

Equating the values given in equations 9.262 and 9.264;

$$(r+x)h(T_L - T_A) = \frac{k}{\ln \frac{r+x}{r}}(T_S - T_L)$$

$$a = (r+x)\frac{h}{k} \ln \frac{r+x}{r} = \frac{T_S - T_L}{T_L - T_A}$$

$$aT_L - aT_A = T_S - T_L$$

$$T_L = \frac{T_S + aT_A}{a+1}$$

Then:

Substituting in equation 9.262:

$$Q = 2\pi l(r + x) \left\{ \left(\frac{T_S + aT_A}{a + 1} - T_A \right) \right\} h$$

$$= \frac{2\pi l h(r+x)}{a+1} (T_s - T_A)$$

$$= 2\pi l h(T_s - T_A) \left\{ (r+x) \frac{1}{1+(r+x)\frac{h}{k} \ln\left(\frac{r+x}{r}\right)} \right\}$$

(9.265)

piferentiating with respect to x:

$$\lim_{t \to 0} \frac{dQ}{dt} = \frac{\left\{ 1 + (r+x)\frac{h}{k} \ln \frac{r+x}{r} - (r+x) \left[\frac{h}{k} \ln \frac{r+x}{r} + (r+x)\frac{h}{k} \frac{r}{(r+x)} \frac{1}{r} \right] \right\}}{\left[1 + (r+x)\frac{h}{k} \ln \left(\frac{r+x}{r} \right) \right]^2}$$

The maximum value of $Q(Q_{\text{max}})$ occurs when dQ/dx=0.

1 -
$$(r + x)\frac{h}{k} = 0$$

 $x = \frac{k}{h} - r$ (9.26)

When the relation between heat loss and lagging thickness exhibits a maximum for the imaged pipe (x = 0), then:

$$\frac{hr}{k} = 1 \tag{9.267}$$

When hr/k > 1, the addition of lagging always reduces the heat loss.

When hr/k < 1, thin layers of lagging increase the heat loss and it is necessary to exceed the critical thickness given by equation 9.266 before any benefit is obtained from the leging. Substituting in equation 9.265 gives the maximum heat loss as:

$$Q_{\text{nax}} = 2\pi l h (T_S - T_A) \left\{ \frac{k}{h} \frac{1}{1 + \frac{k}{h} \frac{h}{hr}} \frac{k}{hr} \right\}$$

$$= 2\pi l (T_S - T_A) k \frac{1}{1 + \ln \frac{k}{hr}}$$
(9.268)

For an unlagged pipe, x=0 and $T_L=T_S$. Substitution in equation 9.262 gives the rate of ball loss Q_s as:

$$Q_o = 2\pi l r (T_S - T_A) h$$
 (9.269)

$$\frac{Q_{\text{max}}}{Q_n} = \frac{k}{rh} / \left(1 + \ln \frac{k}{rh} \right) \tag{9.270}$$

The ratio Q/Q_o is plotted as a function of thickness of lagging (x) in Figure 9.94.

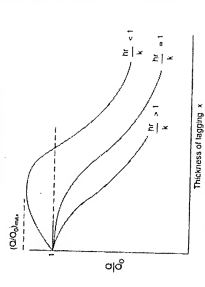


Figure 9,94. Critical thickness of lagging

Example 9.34

A pipeline of 100 mm outside diameter, carrying steam at 420 K, is to be insulated with a lagging manner If the value of heat energy is 7.5 × 10⁻⁴ EMJ and the capital cost of the lagging is to be depreciated and years with an effective simple interest rate of 10 per cent per annum based on the initial investment, when which coass £10/m3 and which has a thermal conductivity of 0.1 W/m K. The ambient temperature may be a as 285 K, and the coefficient of heat transfer from the outside of the lagging to the surroundings as 10 Wird

the economic thickness of the lagging?

Is there any possibility that the heat loss could actually be increased by the application of too thin a the

Solution

For a thick-walled cylinder, the rate of conduction of heat through lagging is given by equation 9.21:

$$Q = \frac{2\pi l k(T_i - T_u)}{\ln(d_o/d_i)} \quad W$$

where d_o and d_i are the external and internal diameters of the lagging and T_o and T_i the corres

Substituting k=0.1 W/mK, $T_o=420$ K (neglecting temperature drop across pipe wall), and $d_i=0.1$, in then

$$\frac{Q}{I} = \frac{2\pi \times 0.1(420 - T_o)}{\ln(d_o/0.1)} \text{ W/m}$$

The term Q/l must also equal the heat loss from the outside of the lagging.

$$\frac{Q}{l} = h_o(I_o - 285)\pi d_o = 10(I_o - 285)\pi d_o \text{ W/m}$$

$$T_{o} = \left\{ \frac{Q}{l} \frac{1}{10\pi d_{o}} + 285 \right\} \quad K$$

$$Q = \frac{2\pi \times 0.1 \left[135 - \frac{Q}{l} \frac{1}{10\pi d_{o}} \right]}{10\pi d_{o}}$$

Thus

HEAT THANSFER

$$\frac{Q}{l} = \frac{2\pi \times 0.1 \times 135}{\ln(d_o/0.1) + 2\pi \times 0.1 \times \frac{1}{107d_o}} = \frac{84.82}{\ln(d_o/0.1) + (0.02/d_o)}$$
W/m

 $_{\rm ac}$ of hear lost = 57.5×10^{-4} /NJ

$$\frac{1}{10} = 84.82 \times 10^{-3} \times 7.5 \times 10^{-4} \times 10^{-6} = \frac{6.36 \times 10^{-3}}{\ln(d_o/2.1) + (0.02/d_o)} \text{ } 2/ms$$

Where of lagging per unit pipe length = $\frac{\pi}{4}[d_n^2 - (0.1)^2]$ m³/m

conditions of largeling = £10/m³ or
$$\frac{\pi}{4}[d_o^2 - 0.01]10 = 7.85(d_o^2 - 0.01)$$
 E/m

Noing that I year = 31.5 Ms, then;

$$t_{\rm merci}$$
 charges = $(0.1 \times 7.85)(d_o^2 - 0.01)/(31.5 \times 10^6) = 2.49 \times 10^{-3}(d_o^2 - 0.01)$ E/Ms

That capital charges = 7.47 ×
$$10^{-3}(d_o^2 - 0.01)$$
 L/Ms

that cost (capital charges + value of heat lost) is given by:

$$C = \left\{ \frac{6.36}{\ln(d_{\rm b}/0.1) + (0.02/d_{\rm b})} + 7.47(d_{\rm b}^2 - 0.01) \right\} 10^{-3} \text{ E/Ms}$$

Differentiating with respect to da:

$$10^3 \frac{dC}{dd_o} = 6.36 \left[\frac{-1}{[\ln(d_o/0.1) + (0.02/d_o)]^2} \right] \left[\frac{1}{d_o} - \frac{0.02}{d_o^2} \right] + 7.47(2d_o)$$

In order to obtain the minimum value of C, dC/ddo must be put equal to zero.

$$\frac{1}{[\ln(d_o/0.1) + (0.02/d_o)]^2} = \frac{(7.47 \times 2)}{6.36} \left[\frac{d_o}{(1/d_o) - (0.02/d_o^2)} \right]$$

$$\frac{1}{[\ln(d_o/0.1) + (0.02/d_o)]^2} = 2.35 \frac{d_o^2}{(d_o - 0.02)}$$

A final and error solution gives $d_0 = 0.426 \mathrm{m}$ or 426 mm. Thus, the economic thickness of lagging = $(426-100)/2 = 163 \mathrm{mm}$

$$hr = \frac{10 \times (50 \times 10^{-3})}{t} = 5$$

For this pipeline:

From equation 9.267, the critical value of hr/k, below which the heat loss may be increased by a thin layer of $\log_{10} k$ is 1. For hr/k > 1, as in this problem, the situation will not arise.

9.12. FURTHER READING

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